#### FINAL REPORT

# CONTRACT NAS2-4310

# STUDY, STERILIZATION AND STORAGE COMPATIBILITY OF GROWTH MEDIA FOR EXTRATERRESTRIAL USE

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#### ABSTRACT

Because of the requirement of dry-heat sterilization for interplanetary spacecraft, an investigation was initiated to assess the scope of the problem presented by the thermal instability of substances important to extraterrestrial life detection experiments. Ninety-four substances were exposed to drastic thermal conditioning. Following heating, thirty-seven of the substances were found to pass preliminary screening tests designed to detect degradation. Twenty of the 57 substances not passing the screening tests were not grossly degraded.

The substances were individually packaged under vacuum in borosilicate glass ampoules. Before packaging most substances were dried in a vacuum. After drying the substances were placed in ampoules which were flushed with nitrogen, evacuated and sealed.

The thermal processing consisted of two separate 92-hour heating periods at  $135^{\circ}$ C. After the second thermal treatment, the specimens were examined by a preassigned sequence of physical and chemical tests designed to detect degradation. Once significant degradation was detected, the test sequence was terminated for that substance.

Candidate substances were selected on the basis of their importance in microbiological growth media, as spore germinating agents, as substrates for metabolic assays and as substances characteristic of major biochemical classes. The forms of the substances tested were chosen on the basis of physical properties and purity of commercially available forms.

Results of tests completed suggest further investigations will find additional evidence of degradation in the heated test substances, but they also suggest that choice of other forms of some of the degraded materials and other packaging conditions will permit them to survive the thermal process used.

#### FOREWORD

This report describes the technical approach and results of an investigation to explore the sterilization and storage compatibility problems of microbiological growth media substances of potential use in the biological exploration of the planets. The investigation was performed by the Space and Re-entry Systems Division of Philco-Ford Corporation under Contract NAS2-4310 with the Ames Research Center of the National Aeronautics and Space Administration. The program was directed by Dr. John B. Opfell of SRS. J. D. Albert, D. E. Gelvin, J. W. Mason, T. A. Oda, and E. R. Walwick of the Philco-Ford, Aeronutronic Applied Research Laboratories performed the laboratory investigations and are the principal authors of this report.

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#### SECTION I

#### INTRODUCTION

This report describes an investigation of the thermal stabilities of substances important to extraterrestrial life detection experiments. The program was a preliminary study designed to assess the problem presented by the inclusion of biological test materials in planetary-impacting spacecraft which require terminal dry-heat sterilization. The thermal conditions chosen for this study were more severe than would be applied in the sterilization of actual planet-bound material. Substances were sealed in glass  $\frac{\text{in }}{92} \frac{\text{vacuo}}{\text{hours}} \text{ and exposed to a temperature of } 135^{\circ}\text{C} \text{ for 2 heating periods of } \frac{\text{vacuo}}{\text{pours}} \text{ and exposed to a temperature of } 135^{\circ}\text{C} \text{ for 2 heating periods of } \frac{\text{vacuo}}{\text{pours}} \text{ and exposed to a temperature of } 135^{\circ}\text{C} \text{ for 2 heating periods of } \frac{\text{vacuo}}{\text{pours}} \text{ and exposed to a temperature of } 135^{\circ}\text{C} \text{ for 2 heating periods of } \frac{\text{vacuo}}{\text{pours}} \text{ and } \frac{\text{vacuo}}{\text{pours}} \text{ and$ 

Candidate substances were selected on the basis of their importance in microbiological growth media, particularly for growth of soil microorganisms, as spore germinating agents, as substrates for metabolic assays and as substances characteristic of major biochemical classes.

Since this was a preliminary investigation detailed descriptions of materials, equipment, and procedures, useful in subsequent investigations are reported.

The literature was reviewed to compile information on the thermal stability of the substances chosen for study. However, little applicable thermal degradation information was found, except for inorganic compounds.

After heating, the thermally treated specimens of each candidate substance were compared with the unheated control specimens by a sequence of both physical and chemical tests. The first group of tests were simple physical tests designed to identify those substance which had been grossly degraded by the thermal treatment. Subsequent tests were more specific and were designed to quantify the degradation. Sufficient time was not available to accomplish this goal in its entirety.

#### SECTION 2

#### SELECTION OF CANDIDATE SUBSTANCES

#### 2.1 LIST OF SUBSTANCES FROM LITERATURE SURVEY

A review of the literature to determine those substances that are particularly useful in microbiological growth media was carried out. Those selected have been used extensively in growth media for soil microorganisms, in classical microbiological growth media, or in spore germinative media. A listing of the substances appears in Appendix A. Excluded from the list were substances used in enrichment media for selective culture.

# 2.2 CHOICE OF CANDIDATE SUBSTANCES

The substances listed in Appendix B were selected as candidates for testing from a compilation of substances used in microbiological growth media (Appendix A), or were included at the request of NASA Ames Research Center. The list of candidate substances was limited to that number which could be procured and handled adequately in terms of the present program. The selection of test substances was based on their: (a) extent of use in general growth media, (b) irreplaceable nature as growth requirements for microorganisms, (c) cost being reasonable in relation to funds available, and (d) estimated physical stability to thermal degradation under the thermal processing conditions. Inorganic salts which were estimated to be stable were not included (see Appendix C, Table I).

Selection of the most useful chemical form of each substance was made after evaluation of physical properties, purity, and cost of available forms. Preferred forms were anhydrous and had high water solubility, high melting point, purity grades of A.C.S., N.R.C., or U.S.P., and low cost.

#### SECTION 3

#### LITERATURE REVIEW

The original purpose of the literature survey was to collect applicable data from laboratory studies on thermal decomposition of the candidate substances. The information obtained was to be used as an aid in constructing test sequences to assess degradation. Even anticipating that little experimental work had been done under similar conditions (135°C and vacuum), it was surprising to find so little information useful to the present effort. The one exception was the data collected on inorganic compounds, which was useful in selecting procedures to detect degradation. The information collected for the organic compounds contributed little to the selection of test procedures. Most procedures which were found would have been proposed on the basis of a fundamental understanding of organic chemistry. The failure of the literature search to uncover much applicable information does serve to justify the need for the present research program.

The literature survey covered the period from 1920 to 1962. Key phrases, in addition to the candidate substances, included: decomposition of, heat of degradation of, heat effects of, thermal stability of. The literature search was limited to the Chemical Abstracts, from which more than 250 papers were located.

Information collected during the search included the identification of products and intermediates formed during pyrolytic, bacterial, oxidative, photolytic, and hydrolytic degradation studies, methods employed in detecting decomposition, and distinctive properties of degraded materials that could be utilized to detect the beginning stages of decomposition. However, most of the data encountered in the literature included decomposition temperatures of compounds, the catalytic effects of contaminants on degradation, thermal studies at a particular temperature and time period,

or the relative stabilities of various compounds at fixed temperatures. Thermal decomposition temperatures could not be used to predict the thermal behavior of candidate substances in the present study unless they greatly exceeded the temperatures of interest,  $135^{\circ}$ C.

#### 3.1 AMINO ACIDS

No thermal degradation studies were found for amino acids, which yielded information concerning degradation products. Photolysis studies (1,2) were encountered that gave evidence indicating the following degradation products were produced: amines with one carbon less than parent amino acids, alcohols, aldehydes,  $CO_2$ , CO, and  $NH_3$ , but not hydroxy- or ketoacids. The photolysis studies could only be used as a rough guide in choosing tests for detection of degradation.

#### 3.2 PROTEINS

Sodium caseinate was reported to be dehydrated below  $153^{\circ}\mathrm{C}$  in 2 hours, and to degrade extensively above  $153^{\circ}\mathrm{C}$  with loss of amine and basic groups (3). At approximately  $300^{\circ}\mathrm{C}$ ,  $\mathrm{CO_2}$ ,  $\mathrm{O_2}$  and  $\mathrm{CO}$  were formed when casein was dry distilled (4). Loss of nitrogen was found to be a function of time and temperature. At  $125^{\circ}\mathrm{C}$  and up to 16 hours, the loss of nitrogen was nearly linear (5). These data were useful in considering tests for caseinate and suggested that changes in solubility and aggregation would be important in evaluating degradation. They would also be useful in considering this protein as a source of amino acids after hydrolysis of the heated caseinate. However, because of the great effort required to determine the amount of each individual amino acid which survived the heating and hydrolysis it was considered expeditious to determine how well the caseinate survived the heating. If it were grossly degraded, further investigation would not be warranted.

#### 3.3 CARBOHYDRATES

Pyrolysis studies of sucrose, glucose, and lactose reported that with temperatures up to  $500^{\circ}\text{C}$  degradation products included  $\text{CO}_2$ , CO,  $\text{C}_n\text{H}_{2n}$ , and  $\text{CH}_4$  and  $\text{H}_2$  (6). These data confirmed that chromatography would be a definitive test for this group of substances and that the screening tests (especially Differential Refractometry) would be appropriate. A chromatographic test for the detection of thermally produced changes in potato starch was described in the literature (7). Accuracy of the test was such that 0.6 mg of decomposed starch in 10 ml of water (0.006% solution) became visible on an alumina column.

#### 3.4 ALCOHOLS AND POLYOLS

Acetaldehyde, hydrogen, and formaldehyde were reported as pyrolysis products of ethanol at  $576-624^{\circ}C$  (8). These results suggested degradation routes that could be evaluated by the test sequences.

#### 3.5 LIPIDS

It was reported that when linoleic acid was heated in a vacuum in the presence of active nickel, the density and the refractive index increased and the iodine value decreased. Without active nickel there was very little change (9). Two of the analysis procedures employed for the tests reported above were included in the test sequence.

# 3.6 VITAMINS

Data on vitamins of even limited value was not encountered in the literature search.

#### 3.7 INORGANIC SALTS

#### 3.7.1 SODIUM THIOSULFATE

The thiosulfates of the alkaline metals, on being heated in the absence of air, are changed into sulfate and polysulfide, and the latter into sulfide and sulfur:  $4\text{Na}_2\text{S}_2\text{O}_3 \rightarrow 3\text{Na}_2\text{SO}_4 + \text{Na}_2\text{S}_5 \rightarrow 3\text{Na}_2\text{SO}_4 + \text{Na}_2\text{S} + 4\text{S}$  (10).

#### 3.7.2 POTASSIUM BICARBONATE

Thermal decomposition of KHCO $_3$  was studied at 140-200°C by means of a thermal balance. The results indicate that decomposition takes place according to KHCO $_3$   $\rightarrow$  5KHCO $_3$  'K $_2$ CO $_3$   $\rightarrow$  2KHCO $_3$  'K $_2$ CO $_3$   $\rightarrow$  KHCO $_3$  'K $_2$ CO $_3$  'K $_2$ C

# 3.7.3 AMMONIUM MOLYBDATE

Differential thermal analysis studies on various forms of ammonium molybdates show that thermal dissociation begins with the removal of water, followed by removal of ammonium hydroxide. The end product in each case is  $MoO_3$  (12).

#### 3.7.4 MANGANOUS SULFATE

Manganous sulfate pentahydrate gave  $\rm MnSO_4\cdot H_2O$  at about  $\rm 300^{O}C$ ,  $\rm MnSO_4$  at  $\rm 800^{O}C$  and finally  $\rm Mn_2O_4$  at  $\rm 900^{O}C$  (13). Manganous sulfate yields MnS at 750-900<sup>O</sup>C, and MnO at 650<sup>O</sup>C on heating 15 minutes with 0-32.5% C in a current of Ar containing not more than 0.03% oxygen (14).

# 3.7.5 FERROUS SULFATE

This salt decomposed completely at  $750^{\circ}$ C (in Ar) to form Fe<sub>2</sub>O<sub>3</sub>, SO<sub>2</sub>, and SO<sub>3</sub> (14).

# 3.7.6 SODIUM PHOSPHATE, DIBASIC

This substance is dehydrated at temperatures above  $240^{\circ}\text{C}$  directly to the metaphosphate with no indication of an intermediate (15).

# 3.7.7 AMMONIUM PHOSPHATE, DIBASIC

This substance decomposes with the loss of 2 moles of NH $_3$  and 2 moles of H $_2$ O. Metaphosphoric acid is always isolated as the semi- or monohydrate (15).

#### 3.8 MISCELLANEOUS

A kinetic study of the decomposition of sodium formate was reported (16). The results indicated two main reactions:

$$2HCOONa \rightarrow Na_2C_2O_4 + H_2$$

$$2HCOONa \rightarrow Na_2CO_3 + H_2 + CO$$

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#### SECTION 4

#### PROCESSING OF CANDIDATE SUBSTANCES

#### 4.1 PROCUREMENT AND STORAGE OF MATERIALS

#### 4.1.1 ALL GLASS CONTAINERS (AMPOULES)

A number of different types of glass containers were tested for durability and ease of handling. The container selected for use was the LG 6770, 10-ml drying ampoule, made by Lab Glass, Inc., Vineland, New Jersey. When received, the ampoules were marked with code numbers, washed, and stored in a covered container until used (refer to Appendix D for the configuration, composition, and preparation of the all glass container.)

#### 4.1.2 CANDIDATE SUBSTANCES

Several sources were considered for each candidate substance. Final choice was based on cost and availability of substances of specified purity. Identification of the procured candidate substances consisted of inspection of the container labels to insure that the material received was that specified. Appendix B gives the source, cost, amount purchased, and manufacturer's lot number for each candidate substance.

Substances were stored in the original, sealed containers, in the dark at room temperature until they were prepared for packaging.

#### 4.2 PREPARATION OF INDIVIDUAL SPECIMENS PRIOR TO THERMAL PROCESSING

# 4.2.1 PREPARATION OF CANDIDATE SUBSTANCES

Since it was considered a disadvantage to have water present in the test

substances, most of them were dried to constant weight in a vacuum oven or vacuum desiccator. Substances were dried at  $80^{\circ}\text{C}$  or at room temperature (23°C). Vacuum desiccators contained  $P_2O_5$  and vacuum pumping was continuous on desiccators and oven during the drying process. Drying conditions for each substance are given in Appendix B. Liquids received no prepackaging treatment.

#### 4.2.2 PACKAGING

Each candidate substance was assigned an identification number. The four replicate specimens of each substance were labeled A, B, C, and D.

Four containers (ampoules) with proper identification numbers were weighed on the Mettler H15 balance. Weights were recorded to the nearest 0.0001 g. A predetermined amount of the prepared test substance was placed in each of the ampoules. Whenever possible approximately 5 g were used. Use of smaller amounts was required when: (a) high cost necessitated purchase of some substances in amounts less than 20 g, (b) 5 g of a substance would have exceeded the capacity of the ampoule, or (c) difficulty was encountered in evacuating ampoules containing finely powdered substances. The weight of one of the heated specimens of each test substance is given in Appendix C, Table I.

Stems of the filled ampoules were heated and drawn out forming a constriction about 3 cm above the shoulder. The constricted section had an inside diameter of approximately 3 mm and a length of 2 cm. They were then connected to a vacuum manifold, evacuated to a pressure of 0.1 mm Hg and returned to atmospheric pressure by filling the manifold with dry nitrogen. After flushing twice with nitrogen the ampoules were evacuated, held at a pressure of 0.1 mm Hg for about 5 minutes, then sealed by melting the glass at the constriction in the stem. The time required to achieve equilibrium with the 0.1 mm Hg vacuum varied with the nature of the test substance. Finely powdered substances presented the greatest difficulty, due to a tendency to escape from the ampoule as the pressure was reduced. In some cases up to an hour was required to reach the 0.1 mm Hg pressure.

Substances not packaged according to the above procedure were ethanol (44), glycerin (45), and beef extract (77). Ampoules containing ethanol were covered with a rubber cap and dipped in liquid nitrogen. When the ethanol was frozen the ampoules were quickly sealed by melting the glass in the stems. Glycerin was packaged in the usual manner except that evacuations were to a pressure of 3 mm Hg instead of 0.1 mm. Beef extract was frozen in liquid nitrogen prior to evacuation of the ampoules.

All specimens were stored in the dark at room temperature.

Pressure measurements during the packaging process were made with a McLeod Vacuum Gauge (Virtis Company, Inc., Gardiner, New York). The gauge and the ampoules being evacuated were attached to the vacuum manifold through stopcocks and rubber sleeves. Distance of the ampoules from the manifold was approximately 15 cm. The most severe constriction in the connecting tubing was the stopcock, with a bore 3 mm in diameter and 17 mm long.

## 4.3 THERMAL PROCESSING AND STORAGE OF PROCESSED SPECIMENS

#### 4.3.1 PROCESSING LOT

Two of the four replicate specimens of each candidate substance were assigned to a processing lot. Each lot consisted of about 40 specimens which were exposed to the thermal process at one time. The other two replicate specimens remained in storage at room temperature. Appendix B shows which of the replicate specimens were exposed to the thermal process and the number of the processing lot to which they were assigned. Five heating lots were used to process the required number of specimens of the 94 candidate substances.

#### 4.3.2 THERMAL PROCESS

The specimens of each processing lot were placed in an aluminum-block tube heater (see Appendix E) which had been stabilized at  $135^{\circ}\text{C}$ . The prescribed 92-hour heat soak period began when the temperature of the specimens reached  $133^{\circ}\text{C}$ . Specimens were held at  $135^{\circ}\text{C} \pm 2^{\circ}\text{C}$  for 92 hours  $\pm$  30 minutes. At the end of the first 92-hour heat soak, the processing lot was transferred to a test tube rack and allowed to cool to room temperature. Specimens were stored in the dark at room temperature between heat soaks. After about 2 days of storage the lot of specimens was placed in the temperature-stabilized heating block for a second heat soak at  $135^{\circ}\text{C} \pm 2^{\circ}\text{C}$  for 92 hours  $\pm$  30 minutes. At the end of the second heat soak the specimens were allowed to cool to room temperature in a test tube rack and were stored in the dark at room temperature.

# 4.3.3 OPENING SPECIMENS FOLLOWING THERMAL PROCESSING

The sealed specimens were weighed before opening to determine if the containers leaked during heating or storage. A possibility existed that containers which had been heated were under pressure resulting from decomposition of test substances. Therefore, all specimens were cooled in liquid nitrogen before opening to condense any decomposition gases which may have been present. A horizontal scratch about 2 mm long was made on each container about 0.5 cm below the shoulder, and a small drop of water applied to the scratch. A hot glass rod pressed firmly against the glass near the scratch caused a crack to form around the container at the level of the scratch. The top of the container was then removed by applying slight pressure with the fingers. After opening, the containers were

covered with a tight fitting rubber cap. During the opening procedure, the specimens were held in a stream of dry nitrogen to minimize entry of air into the opened containers. When the specimens had warmed to room temperature and condensed vapors had dried, the weighing to determine loss of volatiles was performed. Opened specimens were stored in the dark, at room temperature, in screw top jars containing a desiccant.

#### SECTION 5

#### TESTS

#### 5.1 TEST PROTOCOLS

#### 5.1.1 TEST I COLOR CHANGE

Specimens were inspected in their closed ampoules under illumination provided by 16 Sylvania Lifeline No. F30T12-CW-RS fluorescent lamps located in a 9-by-13-foot ceiling which is 6 feet above the inspection station. The color was measured by comparison of the specimen with standard color swatches in the Munsell Book of Color (Glossy Finish Collection)\*. This set of standard swatches represent color in terms of 9 units of value, 18 units of chroma, and 40 units of hue. If the color of the heated specimen differed more than 0.5 units of value, 1 unit of chroma, or 2.5 units of hue from that of the unheated control, the material was considered degraded and no further tests were performed on it.

# 5.1.2 TEST II PHASE CHANGE

Specimens were inspected in their closed ampoules. If either of the following phase changes occurred as a consequence of the heating processes, the material was considered degraded and no further tests were performed on it:

a. The heated material was a liquid whereas the unheated control was a solid, or conversely, the heated material was a solid whereas the unheated control was a liquid.

<sup>\*</sup> Munsell Color Company, Inc., Baltimore, Maryland

b. The heated material was a slush or a solid cake (as distinct from cemented particles) whereas the unheated control was a freely flowing powder or mass of crystals.

#### 5.1.3 TEST III LOSS OF VOLATILES

Specimens producing volatiles were considered degraded and no further tests were performed on them. Specimens were considered to have produced volatiles as a result of the heating processes if they showed one of the following effects:

- a. The ampoule exploded spontaneously during or after the heating process or during the attempt to open it. (The sample was usually lost in the explosion.)
- b. After opening the ampoule the specimen lost more than 0.3% of its weight. Corrections were made for weight changes due to buoyancy and the opening operation (notching).

#### 5.1.4 TEST IV PARTICLES IN SOLUTION

Measured quantities (not more than 50% of the specimen) of the heated specimen and of the unheated control material were each dispersed in enough Millipore-filtered water (Purified Water, U.S.P.) to make solutions which were about 50% saturated. Both solutions were diluted to identical concentrations.

Both solutions were inspected under strong white light against both a flat-black and a flat-white background. If the number of particles visible in the solution from the heated specimen was more than 3 times that in the solution from the unheated control specimen, the material was considered to be degraded and no further tests were performed on it.

# 5.1.5 TEST V pH CHANGE

This protocol was proposed for use in this work but time did not permit its accomplishment.

Measured quantities of the heated specimen and of the unheated control material would be dispersed in enough Millipore-filtered water (Purified Water, U.S.P.) to make 50% saturated solutions. Both solutions would be diluted to identical concentrations and the pH determined either with a pH meter or pH indicator paper. Each solution would be sampled twice with the first sample diluted 50% with water and the second with 0.001 N HCl. If the pH difference between the samples prepared from the heated material differed from that of the samples prepared from the unheated control by more than 1 unit, the material would be considered degraded and no further tests performed on it.

# 5.1.6 TEST VI DIFFERENTIAL REFRACTOMETRY

The solutions prepared for detection of particles in solution (Test IV) served as samples for measurement of differences in index of refraction. The measurements were made in a Brice-Phoenix, Model BP-2000-V, Differential Refractometer. This instrument has a sensitivity in  $\mathbf{n}_D$  of 0.0000013, and the sensitivity of the method is illustrated by the following list of changes in the refractive index of water as solutes are introduced or the temperature of measurement is changed:

Pure water, n<sub>D</sub><sup>20</sup>1.33303 1% Glycerol (aq), n<sub>D</sub><sup>20</sup>1.33416 1% Mannitol (aq), n<sub>D</sub><sup>20</sup>1.33440 1% Dulcitol (aq), n<sub>D</sub><sup>20</sup>1.33441 Pure water, n<sub>D</sub><sup>25</sup>1.33251 2.5% Acetic Acid (aq), n<sub>D</sub><sup>25</sup>1.33427

If the difference between the refractive indexes of solutions prepared from the heated and control specimens was greater than that between two controls, one which is 1% more dilute than the other, the material was considered to have been degraded and no further tests were performed on it.

# 5.1.7 TEST VII X-RAY DIFFRACTION ANALYSIS

The crystal structures of the heated and control specimens were studied with a Philips Norelco X-ray Diffractometer, which used copper  $K\alpha$ -rays. If the difference in the diffraction patterns indicated more than a 1% change between the heated and control specimens the material was considered degraded and no further tests were performed on it.

# 5.1.8 TEST VIII MELTING POINT DETERMINATION

Melting point differences between heated and control specimens were measured with an Electrothermal Melting Point Apparatus\* consisting of an electrically heated block which accommodated three capillary melting point tubes. Simultaneous readings were made on samples of (a) heated specimen, (b) control specimen, and (c) a mixture of heated and control materials.

If there was a difference in melting ranges greater than 3°C, the material was considered degraded and no further tests were performed on it.

<sup>\*</sup> Electrothermal Engineering, Ltd., London, England

#### 5.1.9 TEST IX BARIUM CARBONATE PRECIPITATION

Tests for carbonate ion formation were made on ten-milliliter portions of 1% aqueous solutions of the heated and control specimens. Each solution was treated with six drops of 0.05 Normal barium chloride solution.

If the number of particles visible in the solution of the heated sample was more than twice that in the solution of the control sample, the material was considered to have been degraded and no further tests were performed on it.

#### 5.1.10 TEST X GAS CHROMATOGRAPHY

This protocol was proposed for use in this work, but time did not permit its accomplishment.

Gas chromatography would be performed, either directly on test substances or their solutions, or after conversion of test substances to more volatile trimethylsilyl derivatives. In the case of the amino acids the latter route would be taken:

A sample of the test substance is dissolved in a nonaqueous solvent and treated with N,0-bis-(trimethylsilyl)acetamide (BSA), a silylating agent, and the reaction mixture is injected into the gas chromatograph column. By comparing chromatographic peaks of both control and heated specimens the presence of degradation products can be ascertained.

# 5.2 TEST SEQUENCE

The test sequences indicated in Appendix F for each of the candidate substances were designed principally to detect decomposition products but were not intended to identify these degradation products. Selection of these test sequences was based upon our scientific judgement or, where appropriate, the decomposition studies reported in the literature. However, since each type of test varies in sensitivity, one test in a given sequence may be more conclusive for one substance than another. Consequently, it occasionally could be necessary to complete the entire sequence of tests before conclusions regarding stability of a given substance can be made. Each test was conducted on samples from both control and heated specimens. Direct comparison of the test results for the heated and control specimens gave increased sensitivity of analysis over an attempt to make a judgement of purity of the heated samples alone.

Because the first three tests, for color change, phase change, and loss of volatiles are readily performed and permitted elimination of grossly decomposed substances from further testing, they were performed uniformly on specimens of all candidate substances. Subsequent tests were selectively assigned to various groups of substances. The sequences of these tests are discussed below in paragraphs under the respective headings of these groups.

#### 5.2.1 AMINO ACIDS

The tests for Particles in Solution (IV) and Solution pH (V) were, perhaps, not as sensitive as subsequent tests for members of this series. Because aqueous solutions of these substances were needed in subsequent determinations, gross differences in solubilities between heated and control samples were noted. Buffering capacities, if altered by thermal processing, could also be observed. Differential Refractometry (VI) offered a simple but sensitive procedure to determine if small amounts of degradation had occurred. Gas chromatography offered an alternative sensitive procedure which would permit quantitation of the small amounts of degradation.

#### 5.2.2 PROTEINS

Only the first three tests (I-III) were applied to the sodium caseinate specimens. However, subsequent testing should include amino acid analysis of hydrolyzates for both control and heated specimens. This will determine the appropriateness of proteins to act as a source of amino acids after heating.

Biological assay is also proposed for subsequent work to determine the effectiveness of the heated proteins in supporting growth.

#### 5.2.3 CARBOHYDRATES

a. Monosaccharides. Because neither water-insoluble particles nor acidic products were expected in significant amounts until degradation was quite advanced, the tests for Particles in Solution (IV) and Solution pH (V) were not expected to be particularly sensitive. However, these tests offered simple means to determine intermediate levels of degradation. Solutions of substances used in the tests could also be used for Differential Refractometry (VI). The Differential Refractometry offered a simple but sensitive procedure for detecting changes between heated and control samples, owing partly to the fact that concentrated solutions could be prepared with the sugars. Gas Chromatography (X) (of the trimethylsilylated derivatives) is an established method for determining homogeneity and is included in "Reagent Chemicals", A.C.S. Specifications (1960) for all saccharides listed except salicin, but this substance also is expected to silylate with no difficulty.

b. Oligosaccharides and Polysaccharides. These substances were proposed as sources of monosaccharides and therefore were to be examined after acid hydrolysis. Tests for monosaccharides would be applied to the hydrolyzates.

#### 5.2.4 ALCOHOLS AND POLYOLS

- a. Ethanol and Glycerin. Differential Refractometry (VI) and Gas Chromatography (X) were considered to offer simple and sensitive procedures for the assessment of degradation of glycerin and ethanol.
- b. <u>Mannitol</u> and <u>Dulcitol</u>. These crystalline solids have sharp melting points, therefore, Melting Point Determination (VIII) could be appropriately applied. Gas Chromatography (X) (after trimethylsilylation) would permit quantitation of slight degradation.

#### 5.2.5 LIPIDS

a. Sodium Oleate. This is the salt of an unsaturated fatty acid and can polymerize at high temperatures to form water-insoluble products, therefore, Particles in Solution (IV) was appropriate. Iodine Value Determination could be used to measure the difference in unsaturation between the heated and control samples. This test would be appropriate for detection of intermediate levels of degradation. Gas Chromatography (X) could be performed on the trimethylsilyl derivative to give a very sensitive procedure for detecting degradation. The silylating agent in this case would be trimethylchlorosilane:

 $CH_3(CH_2)_7CH=CH(CH_2)_7COONa + C1 Si(CH_3)_3 \rightarrow CH_3(CH_2)_7CH=CH(CH_2)_7COOSi(CH_3)_3$ 

#### + NaC1

- b. Sodium Acetate. This could be investigated with the Permanganate Reduction to detect decomposition products which would be expected to be one- or two-carbon compounds. Any oxidizable products in the heated sample such as formaldehyde, sodium oxalate, ethanol, etc., would be detected by the consumption of permanganate. This test is included in "Reagent Chemicals", A.C.S. Specifications (1960), p. 437, for sodium acetate. The procedure is to add 5 ml of 10% sulfuric acid and 0.1 ml of 0.1 Normal potassium permanganate to 5 grams of sodium acetate in 50 ml of water. The pink color should not disappear in 1 hour.
- c. <u>Linoleic Acid</u>. This substance is a liquid, unsaturated fatty acid. Differential Refractometry (VI), Iodine Value Determination, Gas Chromatography (X) (of trimethylsilylated derivatives), are all sensitive methods for detection of degradation.

#### 5.2.6 VITAMINS

Thin Layer Chromatography, Ultraviolet Spectroscopy, Fluorimetry, and Melting Point Determination (VIII), are all sensitive tests for the vitamins listed. Particles in Solution (IV) was included as a general exploratory test because many of these substances have complex structures that upon heating would yield water-insoluble resinous degradation products. Differential Refractometry (VI) would be sensitive for those vitamins that are moderately to highly soluble.

#### 5.2.7 PARTIAL HYDROLYSATES

Two separate points to be investigated for these substances are denaturation and amino acid destruction. Particles in Solution (IV) was proposed as an exploratory test to determine aggregation. Amino acid analysis, after complete hydrolysis, would be appropriate for determining whether individual amino acids were destroyed or partially destroyed by the heating. Thin layer chromatography could be used to determine destruction of peptides. Biological assay would be used to demonstrate whether toxic materials were produced by the heating.

#### 5.2.8 NUCLEIC ACID BASES

Many of these substances are only sparingly soluble in water, and solutions for Particles in Solution (IV) would, therefore, be prepared with dilute hydrochloric acid for a comparison of heated with control. These acidified solutions could be used for Thin Layer Chromatography, Differential Refractometry (VI), and Ultraviolet Spectroscopy, all of which are sensitive tests for these compounds.

#### 5.2.9 BIOLOGICAL EXTRACTS

Only biological assays appear to be appropriate additional tests to be performed on these substances.

# 5.2.10 INORGANIC SALTS

a. Sodium Thiosulfate. The literature states that this compound degrades under conditions of heat and no air in the following way:

$$Na_2S_2O_3 \rightarrow Na_2SO_4 + Na_2S + S$$

Particles in Solution (IV) would show any formation of free S. Solution pH (V), is a criterion in "Reagent Chemicals", A.C.S. Specifications (1960). Differential Refractometry (VI), is appropriate because of the solubility of sodium thiosulfate.

- b. Sodium Phosphate, Dibasic, and Ammonium Phosphate, Dibasic. A report in the literature states that these salts convert to a mixture of metaphosphates, PO $_3$ , and pyrophosphates, P $_2$ O $_7$ , at elevated temperatures. X-Ray Diffractometry (VII) is sufficiently sensitive to detect a 0.5% change in the crystal structure.
- c. Potassium Bicarbonate. When heated, this substance is reported to transform entirely to  $K_2CO_3$ . The transformation is stepwise, beginning with the intermediate product 5 KHCO $_3$ · $K_2CO_3$ . Barium Carbonate Precipitation (IX) would detect the formation of any carbonate ions.
- d. Manganous Sulphate, Monohydrate. This salt is reported to form manganous oxide, MnO, and manganous sulfide, MnS, at temperatures above 650°C. If these water-insoluble products are present in the heated sample, Particles in Solution (IV) would detect their presence. Manganous sulfate in the presence of water can also hydrolyze in the following way:

$$Mn(SO_4)_2 + 3 H_2O = H_2MnO_3 + 2 H_2SO_4$$
.

Solution pH would thus also be appropriate.

#### 5.2.11 MISCELLANEOUS

- a. <u>Dipicolinic Acid</u>. Gas Chromatography (X) would be appropriate with the trimethylsilyl derivative.
- b.  $\underline{\text{Furfural}}$ . Differential Refractometry (VI) on the undiluted samples would be quick and conclusive. Gas Chromatography (X) would also give a sensitive test.
- c. <u>Sodium Citrate</u>, <u>Dihydrate</u>. Gas Chromatography (X) would be used on the trimethylsilylated derivative.
- d. <u>Succinic Acid</u>, <u>Lactide and Urea</u>. Melting Point Determination (VII) is simple, sensitive, quick and appropriate for all three compounds.

## 5.2.12 EXOBIOLOGICAL REAGENTS

- a. <u>Sodium Formate</u>. This substance is reported to form sodium oxalate and sodium carbonate. Barium Carbonate Precipitation (IX) would detect any carbonate formed.
- b. <u>Sodium Pyruvate</u>. Gas Chromatography (X) would be used on the trimethylsilyl derivative.
- c. The Naphthylamides (3 substances). Fluorimetry would be performed on these compounds before and after peptidase hydrolysis. To pass this test, fluorescence must not be exhibited by the unhydrolyzed substrate.

#### SECTION 6

#### TEST RESULTS

Test results are given in Appendix G Table I and II. The data indicate that of the 94 candidate substances investigated, 57 failed one or more of the tests. The substances were categorized into one of 5 classes on the basis of the screening test results. The stability class of each substance is given in Table I.

Eleven of the 23 amino acids passed all of the tests with which they were screened. Five of those not passing the tests were only marginally degraded and thus still offer potential, if further effort were expended on finding procedures that would give slight improvement in their stability. No protein, partial hydrolysate, or biological extract passed the screening tests and all were extensively degraded.

Salicin, salicyl alcohol glucoside, was the only survivor of 12 candidate monosaccharides. With the exception of  $\alpha$ -methyl-D-glucoside all of the other monosaccharides were extensively degraded. Of the 5 oligo- and polysaccharides 4 were extensively degraded. Starch survived the heating well and was only slightly discolored.

Three of the four substances in the alcohols and polyols passed all screening tests used. The fourth substance, galactitol, failed because of a slight darkening. However, there was no change in the melting point or mixed melting point (heated + control).

Two of the three lipids passed the screening tests. Linoleic acid, the third substance, was darkened slightly by the heating and failed the color test, but passed all of the other tests used.

Four of the 11 substances in the vitamin group passed all of the screening tests used. Of those not passing, four failed because of slight discolorations or fusing of crystals, but exhibited little deterioration otherwise.

Five of the 8 nucleic acid bases passed the tests imposed and the three that failed did so because of a slight discoloration.

Seven of the nine inorganic substances passed the screening tests used. X-ray diffraction disclosed that the crystal structures of both the heated and control samples of ammonium phosphate (83) were identical, but that the composition corresponded to a mixture of 25 percent  $(NH_4)_2$   $PO_4$  and 75 percent  $(NH_4)$   $H_2PO_4$ .

Three of the substances in the miscellaneous category passed the tests applied. These were dipicolinic acid, succinic acid and sodium formate. Lactide failed because of slight discoloration and urea (m.p. 132) failed because it fused into a cake, sublimed, and lost weight, but remained white in color. None of the amino acid naphthylamides passed, but of these L- $\alpha$ -aspartyl- $\beta$ -naphthylamide seemed least affected by the heating.

#### SECTION 7

#### CONCLUSIONS AND RECOMMENDATIONS

#### 7.1 THERMAL PROCESS SYSTEM

The equipment used in heating and monitoring of heating was found to be completely adequate for the task. If future work called for scaling up the operation, it could best be done by increasing the number of heating and monitoring units rather than increasing the size of the basic heating unit. In initiating this work some consideration was given to the use of an oil bath to achieve the heating process. The oil bath offered two important advantages. Heating of specimens would be more uniform and specimens would be visible during the heating process. Visibility of specimens would permit observation of changes in phase and rate of discoloration during heating. The major disadvantage of the oil bath was the danger presented if an ampoule exploded. Not only would hot oil be projected from the bath, but the exploding ampoule could break other ampoules resulting in loss of specimens. Explosion of ampoules during heating did occur, therefore, this was a real danger. Degradation of the oil in the oil bath also presents a disadvantage.

The drying ampoules used as specimen containers were satisfactory. No changes in ampoule configuration are recommended. The size of the sample of test substance accommodated by the container was found adequate to satisfy the needs of all the tests carried out with a considerable quantity to spare in most cases.

Definite advantages would be gained if an ampoule opening device were constructed to permit evaluation of the pressure developed in ampoules as a result of degradation of test material. Gas samples could be removed from this system and analyzed for degradation components. Packaging

of the test substances presented many problems and consumed a large portion of the experimental effort. The greatest difficulties arose when ampoules were filled with low density powders and with development of static charges on ampoule and substance. It is difficult to transfer the test substance into the ampoule, but even greater difficulties are encountered when the filled ampoule is evacuated prior to sealing. Simple solutions to these problems are not immediately apparent and patience is the most important factor. In future studies it would be an improvement to allow more time for packaging. In those cases where traces of water may decrease stability of the test substances, more time should be allowed on the vacuum manifold prior to sealing. In addition, a vacuum system capable of a better vacuum could be employed.

#### 7.2 EXPERIMENTAL

The amino acids, as a class of compounds, are rather stable to the heating process. However, some amino acids were grossly degraded. Because of the extensive degradation of the proteins, it would not appear reasonable that polymers of individual amino acids will improve their stability. It may be possible to achieve acceptable stability by preparing derivatives of thermally unstable amino acids, even though, results of tests on amino acidnaphthylamides indicate that these derivatives are less stable than the constituent amino acids (Compare results of tests on aspartic acid and phenylalanine with their respective naphthylamides). Another approach that may be value is the investigation of the thermal stability of amino acids in aqueous solution.

The most disappointing group of substances, with respect to thermal stability, were carbohydrates. The stability shown by salicin indicates that the desired stability for monosaccharides can be achieved by making derivatives of them. This is further substantiated by the results obtained with  $\alpha$ -methyl-D-glucoside. The thermal stability demonstrated by starch was also encouraging. This substance was selected for study because it could be used as a source of glucose after acid hydrolysis. Therefore, it can be said that glucose is available for growth media, but some processing (e.g., acid hydrolysis of starch) will be required after heat sterilization of the spacecraft. If the polyols, especially glycerin, are considered satisfactory substitutes for carbohydrates the situation is further improved. Both glycerin and mannitol, and even dulcitol, appear to be good candidates because of their stability to heat. While dulcitol changed color slightly, the lack of difference in melting point when compared with the unheated control, and the lack of change when a mixed melting point was taken, indicates that degradation could not be appreciable.

The vitamins did not fare well as a class, but several of those that failed still hold promise. One example was pimelic acid, although it was discolored by the heating, its crystalline nature was apparent when it solidified (m.p. 103-105°C) upon cooling.

That high decomposition points and melting points of substances are not always good indicators of thermal stability, was demonstrated by the heating of cytosine and orotic acid. It is reported that cytosine has a decomposition point of 320-325°C and orotic acid has a melting point of 345-346°C. Degradation was indicated after heating for both of these substances by a change in color.

Substances in the partial hydrolysate and biological extract groups will require special processing to qualify for use in exobiological-life-detection-experiments. The possibility of finding procedures to stabilize them, as powders, to dry-heat sterilization would seem remote on the basis of their extensive degradation.

The results with inorganic salts showed that, except for molybdate, any of the ions considered, could be furnished. While ammonium molybdate was degraded, molybdate can not be eliminated on the basis of this work. The finding that the ammonium phosphate was 75% (NH<sub>4</sub>)H<sub>2</sub>PO<sub>4</sub> and 25% (NH<sub>4</sub>)2HPO<sub>4</sub> instead of 100% of the latter is attributed to the prepackaging process. As a prepackaging operation the salt was dried in a vacuum oven for 48 hrs at  $80^{\circ}$ C. The change in composition apparently resulted from loss of ammonia. This conclusion was further established by examining the product directly from the manufacturer's container. The results of the investigation showed that the diammonium phosphate was present in roughly three times the quantity of the monoammonium phosphate. Therefore, the drying operation (vacuum oven -  $80^{\circ}$ C) had reversed the ratios of the two ammonium phosphates.

Lactide, in the miscellaneous group of substances, was chosen as a source of lactic acid because of its physical properties. It is formed from lactic acid by heating at 180-220°C in vacuo below 25 mm Hg, has a m.p. of 125°C and a b.p. of 255°C, and hydrolyzes to lactic acid even in cold water. It may be that the discoloration encountered as a result of heating was due to trace impurities. Because of the importance of lactate in intermediary metabolism and the poor thermal stability of sodium pyruvate, further work on improving the stability of lactide should be expended.

#### 7.3 RECOMMENDATIONS FOR FUTURE WORK

#### 7.3.1 GENERAL COMMENTS

This exploratory study placed primary emphasis on physical and chemical detection of thermal degradation of substances typical of microbiological media and representative of biochemical intermediates. Ultimately, the significance of this degradation must be established through suitable tests. Two avenues can be pursued. One would involve the extensive characterization of those compounds which can survive the terminal heat sterilization of the spacecraft and whose contaminants are identified. The other approach is to use the heated materials in the biological tests for which they are intended and determine how well they perform. For example, degradation of growth media components can result in stimulation, inhibition, or no effect

on microbial growth. It can be argued that classification of substances into which are stimulatory and which are inhibitory or inert cannot be made for extraterrestrial organisms because their metabolic requirements and biochemical processes are not known. By the same argument, the classification of substances into metabolites and nonmetabolites cannot be made. The use of growth media sterilized by dry-heat for exobiological studies will be formulated on the metabolic processes and requirements of certain specific terrestrial organisms.

The sum of the degradation products obtained from separate sterilization and storage of a set of growth medium components can differ in both quantity and number compared to degradation products resulting from sterilization of the same components as mixtures. This effect is the result of the increased possibilities for degradation reactions in complex mixtures. an earlier study (under Contract NASw-1065) it was found that degradation products in dry-heat-sterilized complex growth media were generally inhibitory toward growth of microorganisms. This effect was not uniform with respect to all microbial species tested nor was it stable on extended storage. One other useful result of this earlier study was the observation that systems of substances can be fully effective for a particular intended purpose even though each of the separate components suffers degradation in the course of thermal sterilization or storage. As long as the greater part of the specific activity or material survives the sterilization, it is presumably useful in some permutation of the other components of a growth medium. This same concept, of testing systems before developing perfect parts, has been successfully applied in the development of the Saturn V. In successful application it is only necessary to know that none of the parts will fail catastrophically under the conditions of intended usage.

Arguments should also be presented in favor of microbiological media and biochemical test systems composed solely of relatively undegraded constituents. The advantage of such systems is knowledge of all components in the system. This permits manipulation of variables to determine their effect on the system. In the case of degraded materials, lack of knowledge of the composition of the system limits the information derived from changing a variable.

#### 7.3.2 PHYSICAL AND CHEMICAL SCREENING

The first investigation in this sequence will be concerned with the completion of the sequence of physical and chemical tests for those substances so far passing all tests applied to them. This investigation will include a quantitative chemical determination of the extent of degradation of those substances which failed but were not extensively degraded. These two tasks will identify those substances which require little or no further work to meet the requirements imposed by the dry-heat sterilization process. This investigation may also include addition of new candidate substrates.

# 7.3.3 BIOLOGICAL ASSAY SCREENING

The second investigation will be concerned with measurement of the biological usefulness of the thermally processed substances. The investigation will be applied first to the individual substances and then to increasingly complex mixtures of them. The biological usefulness test will be designed to assay the individual, heated, pure substances and mixtures of them for the presence of either growth inhibitory or stimulatory substances for a set of microorganism types representing the spectrum of metabolic systems of greatest interest in exobiological research.

# 7.3.4 DEGRADATION PROCESS STUDY

The third investigation will be concerned with the natures of the degradation process and the products produced by the thermal treatment and storage. A thermodynamic-analysis approach may yield clues about which degradation products to expect. The use of a residual gas analyzer, in concert with a high vacuum chamber in which open containers of the substances are heated, may yield information about the kinetics of the thermal degradation processes.

#### 7.3.5 ALTERNATIVE FORMULATION STUDY

The fourth investigation will be concerned with alternative preparation, formulation, and packaging designs for substances essential for growth media for extraterrestrial use and not among the substances surviving the thermal regime imposed in the current study. Among these alternatives are those which keep separate the reactive groups of the molecules during the high temperature soak, e.g., by solvation, by adsorption on a thermally stable polymer, or by making inert the thermally activated groups.

#### 7.3.6 ALTERNATIVE PROCESSING

The fifth investigation will be concerned with alternative methods for preparing essential growth media substances which are inherently thermolabile. This effort will involve finding other forms of the substances which are more stable and other processes and procedures which will improve the stability of the substances. For each substance, such studies will require extensive expenditure of effort. It is therefore important to establish an order of priority for candidate substances. Those substances considered essential and for which the greatest promise of achieving stability is present, should be given the highest priority. Procedures which may improve stability include: (a) seal substances under hard vacuum, (b) seal substances under pressure supplied by inert gas or gases which result from decomposition of test substances (e.g., ammonia in the ampoule containing ammonium phosphate), (c) process solutions of candidate substances and (e) adsorb test material on a thermostable polymer.

# APPENDIX A

# MICROBIAL GROWTH MEDIA SUBSTANCES

Substance	Reference to Medium Containing Component		
AMINO ACIDS			
β-alanine L-arginine L-arginine L-asparagine L-aspartic acid L-cysteine L-cystine L-glutamic acid L-glutamine glycine L-histidine L-hydroxyproline L-isoleucine L-leucine L-lysine L-methionine L-phenylalanine L-proline L-serine L-threonine L-tryptophan L-tyrosine L-valine	25,26,27 3,12,15,16,18,20,27,33,34,37 7,12,15,20 2a,e,3,5,6,12,15,20,35,37 15,20 1,15,18 6,11,15,20 1,3,7,12,15,20,26,37 15,20,26 15,20 7,15,20 20 7,15,20 9,12,15,20,33 7,15,20 7,10,15,20 10,15,20 12,15,20 12,15,20 12,15,20 12,15,20 12,15,20 12,15,20 12,15,20 12,15,20 15,20 17,12,15,20 17,12,15,20 18,20 18,20 19,15,20 19,15,20 19,15,20 19,15,20 19,15,20 19,15,20 19,15,20 19,15,20 19,15,20 19,15,20 19,15,20		
PEPTIDES			
glutathione	26		
PROTEINS			
casein or sodium caseinate	2,6 21,23		
CARBOHYDRATES			
Monosaccharides D-glucose D-fructose D-galactose	1,2,3,5,6,7,11,12,16,17,20,21,23,26,27,29,30,35,37,38 17,26,35 17,26		

# Monosaccharides (continued)

L-arabinose	26
D-xylose	25,26
D-mannose	17,26
L-rhamnose	26
N-acetylglucosamine	15,17
D-glucosamine	17
$\alpha$ -methyl-D-glucoside	17
2-deoxy-D-glucose	17
salicin	26

# Oligosaccharides

sucrose	3,17,26
maltose	26
lactose	17,26
trehalose	26
melibiose	26
melezitose	26
gentianose	26
raffinose	26

# <u>Pol</u>ysaccharides

starch	26
inulin	26
cellulose	26
dextrin	26
glycogen	26

# ALCOHOLS AND POLYOLS

ethanol	16,29
glycerol	3,25,26
D-mannitol	2a,5,6,17,22,24,25,26
D-sorbitol	26
D-galactitol	26
adonitol	26

# LIPIDS

oleic acid or sodium salt	9
acetic acid or sodium salt	3,7,26
linolenic acid	15
linoleic acid	9
arachidonic acid	9
mevalonic acid	15
cholesterol	15
lactobacillic acid	15

# LIPIDS (continued)

vaccinic acid 15 9- or 10-hydroxy oleate 15 9- or 10-hydroxy stearate 15

#### VITAMINS

D- or (+)- biotin 1,3,7,15,25,26 thiamine 1,3,7,15,20,25,26 thiamine pyrophosphate 3,15,20,26,29 nicotinic acid 15,26 nicotinamide riboflavin 3,15,20,26 flavin mononucleotide 15 flavin adenine dinucleotide 15 3,15 pyridoxine -15,26 pyridoxal 3,15,20 pyridoxamine pyridoxine phosphates 15 para-aminobenzoic acid 3,15,20,25,26 folic acid 3,15,20,26 1,15,26 cobalamin myo-inositol (L-) 3,15,26 3,15,20,26 choline pantothenic acid 3,15,20,26 15 pantetheine 15 lipoic acid 15 vitamin K iron porphyrins 15,25 26 hematin heme ferrichrome coprogen 26 pimelic acid 15 terregens factor

## PARTIAL HYDROLYSATES

6,8,11 casitone 3,5,14,22,23 peptone 6d, 14, 31 soypeptone 6d,7,14,31 tryptone 14,38 tryptose 38 proteose peptone #3 12 casamino acids 5,14 multi-peptone

# NUCLEIC ACID BASES

3,15,20,26
3,15,20,26
15
3,15,20,26,29
20,26
15,26
20,26
3,15,20,26

# NUCLEOSIDES

adenosine	15,27
adenine deoxyriboside	15
guanosine	15
guanine deoxyriboside	15
inosine	
hypoxanthine deoxyriboside	10, 15, 2

hypoxanthine deoxyriboside 10,15,27 xanthosine 15 cytidine 15 cytosine deoxyriboside 15 thymidine 15 uridine 15

# NUCLEOTIDES

adenylic acid	26
adenosine-5'-phosphate	15
guanylic acid	15,26
inosine-5'-phosphate	15
cytidylic acid	15
uridine-5'-phosphate	15

# NUCLEIC ACIDS

yeast RNA 20

# BIOLOGICAL EXTRACTS

yeast	3,6d,e,8,11,28,29,36
beef	5,14,23,28
beef heart; infusion	8,14,38
malt	3
calf brain; infusion	14,38
soil	2,6,23

#### INORGANIC SALTS

```
sodium chloride
                                    2,3,5,6,20,22,24,31,38
   carbonate
   nitrite
                                    4,13,29
   nitrate
                                    3,8,29
   molybate
                                    3,20,22,29
   sulfate
                                    1,3,6,12,20,29
   sulfite
                                    14
   thiosulfate
                                    29
   monohydrogen phosphate
                                    6,14,37,38
   dihydrogen phosphate
                                    6,20,37
   bicarbonate
                                    1
   silicate
                                    3
   pyrophosphate
                                    34
   sulfide
                                    3,29
                                    3,20
potassium chloride
   nitrite
                                    1,4
   nitrate
                                    2,5,23
   monohydrogen phosphate
                                    1,3,4,5,6,8,14,20,22,23,24,29,37
   dihydrogen phosphate
                                    1,3,4,6,12,22,29,30,37
   sulfate
                                    22,24
   bicarbonate
                                    1
ammonium nitrate
                                    3,12,29,30
   chloride
                                    1,3,6,12,20,29,37
   monohydrogen phosphate
                                    3,6
   paramolybdate
                                    3
   sulfate
                                    1,20
calcium chloride
                                    1,2,3,5,6,12,20,29,37
   carbonate
                                    1,20,23,29
   sulfate
                                    2,6
magnesium carbonate
   sulfate
                                    1,2,3,5,6,20,22,23,24,29,30
   chloride
                                    6,12,20,37
boric acid
                                    3,20
ferrous chloride
                                    12
   sulfate
                                    1,3,6,18,29
ferric chloride
                                    1,2,3,5,6,20,22
manganous chloride
                                    3,12,29
   sulfate
                                    1,3,20,22,37
cobaltous nitrate
                                    3
   sulfate
                                    20
cupric sulfate
                                    1,3,20
zinc sulfate
                                    1,3,20
   chloride
                                    3
```

# MISCELLANEOUS

dipicolinic acid furfural	19 32
lactic acid or sodium salt	2,6,15,26,29
sodium citrate	3,20,26
2-furfuryl diacetate	32
2-furfuryl-n-butyrate	32
sodium malate	3,26,29
succinic acid or sodium salt	3,26
oxalic acid	26
betaine	26
ammonium tartrate	3
putrescine	3,26
spermidine or spermine	15
Tris(hydroxymethylamino)-methane	48
λ-pyran-2,6-dipicolinate	19
urea	29

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  - i. Jensen's streptomycete medium, 30 b.
  - j. Starkey's sulfur oxidation, 30 c.
  - k. Beijerinck's thiosulfate oxidation medium, 30 d.
  - 1. Van Delden's sulfate reduction medium, 30 e.
  - m. Ammonification medium, 30 f.
  - n. Heterotrophic iron oxidation, 30 g.
  - o. Burk's nitrogen fixation medium, 30 h.
  - p. Modified carbon nutrition agar, 31 a.
  - q. Leathen's autotrophic iron oxidation, 31 b.
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Peptone + trypticase soy agar

Peptone + tryptone

Peptone + soy peptone, bacteriological

Peptone + NaCl

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Media for growing facultative anaerobes Heart infusion broth (Difco) Brain-heart infusion broth (Difco)

APPENDIX B

CANDIDATE SUBSTANCES; PROCUREMENT, PREPARATION AND THERMAL PROCESSING DATA

Substance and Quality*	Empirical Formula	Wgt.		aninos	Lot No.	dollars/gram Recd. Opened Pdgd. Lot Specimen	DEC.		1			Prepackaging drying	(4× 8)
AMINO ACIDS													OC hrs
β-Alanine, Μ.A.	$c_3 H_7 NO_2$	89.1	1	MRI	\$2321	3.50/100	9-20	9-25	9-28	٣	A,B	vac oven	23 48
L-q-Alenine, N.R.C.	$c_3 H_7 NO_2$	89.1	2	CBI	57251	12.50/50	9-13	9-25	9-28	3	А,В	vac oven	23 48
L-Arginine hydrochloride, N.R.C.	$C_{6}H_{14}N_{4}O_{2}\cdot HC1$	210.7	3	CBI	59323	3.25/50	9-13	9-14	9-18	1	B, D	vac oven	23 20
L-Asparagine, N.R.C.	C4H8N2O3	132.1	4	CBI	59419	4.00/50	9-13	9-25	9-29	33	A,B	vac oven	23 48
L-Aspartic acid, N.R.C.	C4 H7 NO4	133.1	2	GBI	59859	3.50/50	9-13	9-14	9-22	2	c,b	vac oven	80 20
L-Cysteine hydrochloride monohydrate, N.R.C.	$c_3 + 1 + 10 + 10 + 10 + 10 = 10$	175.8	9	189	80193	3.25/50	9-13	9-14	9-22	2	A,B	vac oven	80 40
L-Cystine dihydrochloride	$c_{6}{}^{H_{12}}{}^{N_2}{}^{O_4}{}^{S_2} \cdot {}^{2}$ HC1	313.2	7	CBI	808	6.50/50	9-13	9-14	9-18	-	A,B	vac oven	80 20
L-Glutamic acid hydrochloride, C.P.	$c_5 H_9 NO_4 \cdot HC1$	183.6	00	MRL	R2965	4.50/100	9-20	9-25	9-29	3	А,В	vac oven	23 48
L-Glutamine, N.R.C.	$c_{5}H_{10}N_{2}^{0}$	146.6	6	GBI	58142	11.00/50	9-13	10-10	10-16	5	А, В	vac oven	23 144
Glycine, N.R.C.	$c_2^{H_5NO_2}$	75.1	10	CBI	59264	1.25/50	9-13	9-25	9-29	ъ	А,В	vac oven	23 48
L-Histidine, N.R.C.	C6H9N3O2	155.2	11	CBI	59184	10.00/50	9-13	9-25	9-29	9	А,В	vac oven	23 48
4-Hydroxy-L-proline, N.R.C.	с <sub>5 Н9</sub> NO <sub>3</sub>	131.1	12	GRI	44452	18.75/25	9-13	9-25	9-29	3	А,В	vac oven	23 48
L-Isoleucine, N.R.C. (allo free)	C <sub>6</sub> H <sub>13</sub> NO <sub>2</sub>	131.2	13	GBI	80717	21.25/25	9~13	9-14	10-9	4	А,В	vac desicc	23 48
L-Leucine (methioning free)	C <sub>6</sub> H <sub>13</sub> NO <sub>2</sub>	131.2	14	GBI	95965	4.75/50	9-13	9-25	9-29	Э	А,В	vac oven	23 48
L-Lysine hydrochloride, N.R.C.	C <sub>6</sub> H <sub>14</sub> N <sub>2</sub> O <sub>2</sub> ·HC1	182.7	15	GBI	80745	3,25/50	9-13	9-14	9-22	2	c,b	vac oven	80 20
L-Methionine, N.R.C.	$C_5H_{11}NO_2S$	149.2	16	GBI	82532	8.00/50	9-13	9-25	9-28	3	A,B	vac oven	23 48
L-Phenylalanine, N.R.C.	$c_9 H_{11} NO_2$	165.2	17	GBI	57104	7.50/25	9-13	9-25	10-2	3	A, B	vac oven	23 48
L-Proline, N.R.C. (hydroxy-L-proline free)	$c_5 H_9 NO_2$	115.1	18	GBI	651421	8.75/25	9-13	9-14	10-20	4	A,D	vac desicc	23 48
L-Serine, N.R.C.	C3H7NO3	105.1	19	GBI	58270	15.00/25	9-13	9-18	9-22	7	c,D	vac oven	80 20
L-Threonine, N.R.C. (allo free)	C4 H9 NO3	119.1	20	CBI	59657	15,00/25	9-13	10-3	10-6	ς.	А, В	vac oven	23 48
L-Tryptophan, N.R.C.	$c_{11}^{H_{12}^{N_2}O_2}$	204.2	21	GBI	59858	14.50/25	9-13	10-3	10-19	5	C,D	vac oven	23 48
L-Tyrosine, N.R.C.	$c_{9}$ $H_{11}$ $NO_{3}$	181.2	22	CBI	80819	3,25/50	9-13	10-3	10-6	2	А,В	vac oven	23 48
L-Valine, N.R.C.	$c_{5}$ $H_{11}$ $^{NO}$ $_{2}$	117.2	23	GBI	80719	13.00/50	9-13	9-22	9-27	e E	A, B	vac oven	23 72
PROTETH													
Sodium caseinate (soluble casein)			57	CBI	82243	.75/50	9-13	10-3	10-16	S	А,В	vac oven	23 192
CARBOHYDRATES													
Monosaccharides													
N-Acety1.Q-D-glucosamine	C8HSNO6	211.1	25	ACC	022461	16.00/50	9-20	10-10	10-16	2	в, D	vac oven	23 48
L-Arabinose, N.R.C.	C5H10O5	150.1	27	GBI	82582	05/00.9	9-13	9-14	9-21	2	Q	vac oven	80 20
2-Deoxy-D-glucose, N.R.C.	$c_{6}^{H_{12}^{O}}$ 5	164.2	28	CBI	82975	36.25/25	9-28	10-3	10-11	4	А,В	vac oven	23 48
Levulose (B-D-fructose), N.R.C.	C6H12O6	180.2	29	GBI	81043	1.50/50	9-13	9-14	9-27	m.	A,B	vac oven	23 72
D-Galactose, N.R.C.	C <sub>6</sub> H <sub>12</sub> O <sub>6</sub>	180.2	30	GBI	81030	3,00/50	9-13	9-14	9-22	7	υ	vac oven	80 20
D-Glucosamine hydrochloride	C <sub>6</sub> H <sub>13</sub> NO <sub>5</sub> ·HC1	179.2	31	GBI	29004	4.00/50	9-13	9-14	9-21	7	C,D	vac oven	80 20
Dextrose ( $lpha ext{-D-glucose}$ ), reagent, A.C.S.	C <sub>6</sub> H <sub>12</sub> O <sub>6</sub>	180.2	32	B&A	1602	1.96/227	9-11	9-18	9-27	m	A,B	vac oven	23 72
D-Mannose, N.R.C.	C <sub>6</sub> H <sub>12</sub> O <sub>6</sub>	180.2	33	CBI	57126	5.00/50	9-13	9-18	9-27	m	A, B	vac oven	23 72
α-Methyl-D-glucoside	$c_{7}^{H_{14}^{0}6}$	194.2	34	MRL	11595	3,50/100	9-20	10-3	10-10	S	А,В	vac oven	23 48
L(+)Rhamnose monohydrate, M.A.	$c_{6}^{H_{12}^{O_5} \cdot H_2^{O}}$	182.2	35	MRL	83918	13.00/50	9-20	10-3	10-6	ν.	А,В	vac oven	23 48
D-Salicin (Bact.), Puriss	C <sub>13</sub> H <sub>18</sub> O <sub>7</sub>	286.3	36	PCC	379100	24.00/50	10-4	10-10	10-16	5	A, B	vac oven	23 48
D-Ribose, N.R.C.	C <sub>5</sub> H <sub>10</sub> O <sub>5</sub>	150.1	37	GBI	81545	18,00/50	9-28	10-3	10-11	4	А,В	vac oven	23 48
Oligosaccharides													
lpha-Lactose monohydrate, U.S.P.	$c_{12}{}^{H_{22}}{}^{0}{}_{11}$ . $^{H_{2}}{}^{0}$	342.3	38	SIC	363-1830	1.00/100	8-31	6-5	9-15	-	A,B	vac oven	23 120
8-D-Maltose monohydrate, N.R.C.	C <sub>12</sub> H <sub>22</sub> O <sub>11</sub> ·H <sub>2</sub> O	360.3	39	GBI	56330	1.75/50	9-13	9-14	9-01	2	A,B	vac oven	23 96

Polysaccharides				
Starch, soluble powder, reagent, A.C.S.			41	B&A
Inulia	(C <sub>6</sub> H <sub>10</sub> O <sub>5</sub> )n	200	7.5	C3C
ALCOHOLS AND POLYOLS				
Dulcitol (D-galactitol), N.R.C.	C6H1406	182.2	4,3	CBI
Ethyl alcohol, pure (ethanol), U.S.P.	c <sub>2</sub> H <sub>6</sub> o	46.1	77	ISI
Glycerine (glycerol), reagent, A.C.S.	c <sub>3</sub> H <sub>8</sub> o <sub>3</sub>	92.1	45	B&A
D-Mannitol, N.R.C.	°6 <sup>4</sup> 1 <sup>4</sup> 0 <sup>6</sup>	182.2	46	GBI
LIPIDS AND RELATED SUBSTANCES				
Sodium acetate, reagent	$c_2 H_3 o_2 Na$	82.0	41	B&A
Linoleic acid, Puriss (99%)	$c_{18}^{H_{32}}c_{2}$	280.4	84	PCC
Oleic acid sodium salt (sodium oleate), practical	C <sub>18</sub> H <sub>33</sub> O <sub>2</sub> Na	304.4	20	¥CB
VITAMINS				
Para-aminobenzoic acid, U.S.P.	$c_7 H_7 NO_2$	137.1	51	GB CB
Biotin, crystalline	$c_{10}^{H_{16}^{N_2}0_3}$	244.3	52	GBJ
Choline chloride, crystalline (99%)	C5H14C1NO	139.6	53	SIC

48

80

vac oven

1 A, F

9-6

9-5

8-4

1.37/454

Z118

B&A

40

342.3

 $c_{12}^{H_{22}^{O}_{11}}$ 

D(+)Sucrose, reagent, A.C.S.

120

23 23

vac desicc vac oven

A,D

S

10-5

9--5

**9-**%

3.00/227

Z133

10-18 10-18 10-22

2.50/25

63156

20

vac oven

9-21

9-14

9-13 9-11

19.50/50

57085

9-14

36.03/2986

53298

20

80

vac oven

9-23

9-18

9-13

1.50/50

80960

7-8

2.59/596

2215

84

80

vac oven

4

10-25

10-4

17.50/25

0352920

9-5

8-31

2.85/1000

0X1X0

9-5

8-31

2,47/454

X113

84

80

vac oven

80

vac oven

9-25

9-18

1.50/50 120.00/10

81885

10-3

9-13

82433

9-5

8-31

.90/100

LIPIDS AND RESORTUM ace Linoletc a Cletc acid VITAMINS Pata-aminc	., (,
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LIP	VIT	

LIPID	Soc	Li	01	VITAM	Pa	B1	ŧ

	LIPIDS A	Sodium	Linole	Oleic	VITAMINS	Para-	Bioti	;
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LIPIDS AN	Sodium	Linole	Oleic	VITAMINS	Para-e	Biotir	tho 147

COT 1111	Sodiu	Linol	Oleic	VITAMINS	Para-	Bioti	Cho1i

Sodium	Linolei	Oleic 6	VITAMINS	Para-an	Biotin	Choline

Para- Bioti Choli	
Biot	
Para-	
VITAMIN	
Oleic	
Linol	
2001	

Linoleic acid, Pur Oleic acid sodium	VIIAMINS Para-aminobenzoic	Biotin, crystallin	Choline chloride,	i-Inositol (meso)
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Sodi	Line	016	VITAMI	Par	Bio	Cho	

Lino Olei VITAMIN Para Biot	VITAMIN Para Bioc	VITAMIN	01ei	Lino	Sodi	
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 Nia	Ca]

120

80 23 23

vac oven

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1.50/50

9-5

2.50/100

71794

123.1

C<sub>6</sub>H<sub>12</sub>O<sub>6</sub>  $c_6 H_5 NO_2$ 

vac oven

120

87

vac oven vac oven

9-21

9-5

8-31

4.20/50

77B-0300

376.4 337.3

 $c_8 H_{11} NO_3 \cdot HC1$ 

C17H20N406

9-20 9-13

7.00/50

53393

ÆĽ

C12H17C1N4OS·HC1

3.50/50

57910

GBI

62

Menadione sodium bisulfite (water soluble Vitamin K)  $c_{11}$   $^{1}$   $^{1}$   $^{8}$   $^{2}$   $^{2}$   $^{1}$   $^{2}$   $^{9}$   $^{2}$   $^{3}$   $^{1}$   $^{2}$   $^{2}$   $^{3}$   $^{1}$   $^{2}$   $^{3}$   $^{3}$   $^{2}$   $^{3}$   $^{2}$   $^{3}$   $^{2}$   $^{3}$   $^{2}$   $^{3}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^{2}$   $^$ 

Casitone\*\* vitamin free, dehydrated

PARTIAL HYDROLYSATES

Proteose Peptone No. 3\*\*

10-3

9-5

4.50/50

105B-1660

10-3

15.00/50

105B-1030

16.00/50

12196

160.2 205.6

476.5

 $(c_9H_{16}N_5)_2$ ca

none

10-13 10-13 10-13 10-13 10-18

10-4 10-4 10-4

3.50/114 2.55/114 2,20/114

496024

064964

510127

DIF

99

'n

10-18

vac oven

9-25

9-18

9-13

13.50/50

80460

GBI

135.1

C<sub>5</sub>H<sub>5</sub>N<sub>5</sub>

vac oven vac oven vac oven

10-24 4 A,B

9-5

24.50/50

24.50/25

81727

9-25

9-22 9-18

9-13 9-13 8-31

17.00/50

GBI GBI

151.1 136.1 156.1

 $c_5 H_5 N_5 0$ C4H2N3O

C2H4N40  $C_5H_4^4N_2^04$ 

Orotic acid (uracil 4-carboxylic acid)

Hypoxanthine, N.R.C.

Cytosine, N.R.C.

Guanine

NUCLEIC ACID BASES Bacto-soytone\*\*

Adenine, N.R.C.

11111

9-13

38,00/25

NIACI	Calci	

Calcium-D-pantothenate, U.S.P.	Pimelic acid, C.P.	Pyridoxine monohydrochloride (Vitamin B6)	Riboflavin (Vitamin B2)	Thiamine hydrochloride, (Vitamin Bl), U.S.P.
--------------------------------	--------------------	-------------------------------------------	-------------------------	----------------------------------------------

in (nicotinic acid) U.S.P.

יחדום דחר הוו חווים ביודוול וודים ביודוול וודים ביודוול וודים	2,6,1,9,1,5	11091	ţ			W. 11.1.170		5.	2 23	-				
Uracil	C, H, N, O,	112.1	7.5	GBI	59551	5,00/50	9-13	10-10	10-17	4	A,B	vac oven	23 48	
Xanthine	C5H4N4O2	152.1	92	NBC	1963	13.38/50	8-31	9-6	9-22	2	B,D	vac oven	80 48	
BIOLOGICAL EXTRACTS														
Beef extract			7.7	DIF	493821	5.25/114	10-4	10-30	10-30	4	C,D	none		
Beef heart for infusion			78	DIF	511085	13,00/454	10-4	10-27	10-28	7	A,B	none		
Malt extract			62	DIF	491155	1.05/114	10-4	10-18	10-18	4	A,B	none		
Yeast extract			80	DIF	493665	3.70/114	10-4	10-25	10-25	4	A,B	none		
INORGANIC SALTS														
Armonium chloride, granular, reagent, A.C.S.	NH, C1	53.5	81	B&A	A086	1.55/454	8-4	9-2	9-12	1	А,В	vac oven	23 120	
Ammonium molybdate, crystal, reagent, A.C.S.	$(NH_4)_6 Mo_7 0_{24} \cdot ^4H_2 0$	1236.0	82	B&A	Y364	2,46/114	8-31	9-5	9-15	1	А,В	vac oven	80 48	
Ammonium phosphate, dibasic, crystal, reagent, A.C.S.		132.1	83	B&A	2040	2,46/454	8-4	9-5	9-15	1	A,B	vac oven	87 08	
Ferrous chloride, crystal, reagent		126.8	78	B&A	Y309	2.09/114	8-31	10-18	10-19	4	A,B	vac desicc	23 48	
Ferrous sulfate, crystal, reagent A.C.S.	Feso, 7H20	278.0	85	B&A	A025	1.82/454	8-4	9-15	9-23	2	C,D	vac desicc	23 168	
Manganous sulfate monohydrate, powder, reagent, A.C.S. MnSO, H,O	O'H-7OSUM.	169.0	98	JTB	34454	2,98/227	8-4	9-5	9-21	2	C,D	vac oven	87 08	
Potassium bicarbonate, crystal, reagent, A.C.S.	кнсо3	100.1	87	JTB	34035	1.96/454	7-8	9-5	9-12	1	A,B	vac oven	80 48	
Sodium phosphate, dibasic, reagent, A.C.S.	Na <sub>2</sub> HPO <sub>4</sub>	142.0	88	B&A	X366	2.74/227	9-8	10-10	10-18	2	А,В	vac oven		
Sodium thiosulfate, reagent	$Na_2S_2O_3$	158.1	68	B&A	2138	1,98/454	8-31	9-2	9-14	1	А,В	vac oven	80 48	
MISCELLANEOUS														
Dipicolinic acid (2,6-pyridine dicarboxylic acid)	C7HSNO4	167.1	90	ACC	022051	10,00/100	9-20	10-10		2	A,B	vac oven	23 48	
Furfural, reagent	C <sub>5</sub> H <sub>4</sub> O <sub>2</sub>	96.1	91	B&A	2235	2.84/526	8-31	9-16	9-16	-	A,B	none		
Lactide, reagent	708H <sup>9</sup> 2	144.1	92	K&K	2031	42.00/30	10-4	10-19	10-24	7	A,B	vac desicc		
Sodium citrate dihydrate, crystal, reagent	Na3C6H507.2H20	294.1	93	B&A	X126	1.52/114	8-31	9-5	9-11	1	A,B	vac oven	87 08	
Succinic acid, crystal, reagent	C4H6O4	118.1	76	B&A	2116	2.32/114	8-31	9-2	9-25	2	C,D	vac oven		
L-c-Aspartyl-B-naphthylamide, M.A.	C14H14N2O	258.3	95	MRL	S2881	30.00/1	9-20	9-28	9-29	m	A,B	vac desicc		
L-Histidyl-8-naphthylamide, M.A.	C16H16N40	280.3	96	MRL	\$1257	30.00/1	9-20	9-28	9-29	3	A,B	vac desicc		
L-Phenylalanyl-8-naphthylamide, M.A.	$c_{19}{}^{H_{18}}{}^{N_2}{}^{O}$	290.4	46	MRL	84611	30.00/1	9-18	9-28	9-29	3	A,B	vac desicc		
Sodium formate, crystal, reagent	NaCHO <sub>2</sub>	68.0	86	B&A	2278	4.20/454	8-31	9-5	9-15	1	А,В	vac oven		
Sodium pyruvate, reagent	$c_3 H_3 O_3 Na$	110.0	66	GBI	80550	20.00/50	9-13	9-14	9-01	2	A,B	vac oven		
Urea, crystal, reagent, A.C.S.	(NH <sub>2</sub> ) <sub>2</sub> CO	60.1	100	B&A	Y318	2.72/454	8-31	9-5	9-11	1	A,B	vac oven	87 08	
* Except where indicated by A.C.S., N.R.C., or U.S.P., is unique for the manufacturer.	or U.S.P., the quality specification	ation												

B-1-3

\*\* Trademark

## ABBREVIATIONS

A.C.S.	American Chemical Society Specification
U.S.P.	United States Pharmacopeia Specification
N.R.C.	National Research Council Specification
MRL	Mann Research Laboratories, Inc., New York, N. Y.
GBI	General Biochemicals, Chagrin Falls, Ohio
В&А	Baker and Adamson, General Chemicals Division, Allied Chemical Corporation, Morristown, New Jersey
ACC	Aldrich Chemical Company, Inc., Milwaukee, Wisconsin
PCC	Pierce Chemical Company, Rockford, Illinois
JTB	J. T. Baker Chemical Company, Phillipsburg, New Jersey
USI	U. S. Industrial Chemicals Co., New York, N.Y.
CBC	California Corporation for Biochemical Research (Calbiochem), Los Angeles, California
МСВ	Matheson Coleman & Bell Division, The Matheson Company, Inc., East Rutherford, New Jersey
NBC	Nutritional Biochemicals Corporation, Cleveland, Ohio
DIF	Difco Laboratories, Detroit, Michigan
SIG	Sigma Chemical Company, St. Louis, Missouri
K&K	K&K Laboratories, Inc., Plainview, New York
M.A.	Mann Assayed
C.P.	Chemically Pure

## APPENDIX C

## THERMAL STABILITIES OF SELECTED CANDIDATE INORGANIC SALTS

Thermal stability of selected candidate inorganic salts was estimated on the basis of either their low vapor pressures at  $135^{\circ}C$  and 20  $\mu$  Hg or their melting point, or boiling point, or decomposition point being in excess of  $135^{\circ}C$ . The vapor pressures were calculated from thermodynamic free energies of formation of potential gaseous decomposition products at equilibrium with the conditions specified above. The purpose of this evaluation was to identify those inorganic salts which would readily withstand sterilization conditions, and could therefore be eliminated from further testing. Those substances with questionable stability would be selected for testing on the basis of their priority for use in microbiological growth media.

Those compounds judged to be stable were investigated for availability. The ones that were found to be commercially available in the anhydrous or stable hydrated form and in reagent grade or meeting A.C.S. specifications were eliminated from testing. The compounds falling in this category are listed in Table I. The remaining compounds which required laboratory experimentation to establish their ability to meet the heating requirements are listed in Table II.

TABLE I

INORGANIC SALTS - ESTIMATED TO BE STABLE AT 135°C, 20 µ Hg ABSOLUTE PRESSURE

Calculated

		D, dm	Vapor Pressure L Hg (135°C)	b, oc
calcium chloride	$\mathtt{CaC1}_2$	772	•	>1600
carbonate	$caco_3$	d.825	10-10	
cobaltous sulfate	CoSO	686		
cupric sulfate	CuSO <sub>4</sub>	200		d.650
ferric chloride	${ m FeC1}_3$	282		315
magnesium carbonate	$MgCO_3$	d.350		-co <sub>2</sub> ,900
sulfate	$M_{\rm gSO_4}$	d.1124		
potassium chloride	KC1	776		sub1.1500
nitrite	KNO <sub>2</sub>	d.350		
nitrate	KNO <sub>3</sub>	334		d.400
sulfate	$K_2SO_4$	588	Ç	
sodium chloride	NaCl	801	10-12	1413
nitrite	$naNO_2$	271		d.320
nitrate	$\frac{1}{100}$	307	Č	d.380
sulfite	$Na_2SO_3$		10-30	d.
sulfate	$Na_2SO_4$	884		
sulfide	Na <sub>2</sub> S	1180		
carbonate	$Na_2CO_3$	851		ф.
pyrophosphate	$^{\mathrm{Na_4P_2O_7}}$	880		
zinc chloride	$\operatorname{ZnC1}_2$	262		732
sulfate	ZnS0 <sub>4</sub>	d.740		

TABLE II

INORGANIC SALTS REQUIRING LABORATORY DATA TO ESTABLISH STABILITY AT 135°C, 20 µ Hg ABSOLUTE PRESSURE

		D°, qm	Calculated Vapor Pressure L Hg (135 C)	bp, °c
ammonium chloride	$^{ m NH}_{ m c}$ C1	sub1.335	21	
paramolybdate	$(NH_4)_6 Mo_7 O_{24} \cdot 4 H_2 O$	<b>q</b> .		
monohydrogen phosphate	$(NH_4)_2^{2}$ HPO $_4$	d.		
nitrate	NH <sup>4</sup> NO <sup>3</sup>	170		d.210
sulfate	$(NH_4)_2^SO_4$			
boric acid	$^{\mathrm{H}_3\mathrm{BO}_3}$	d.185		$-1^{1/2}$ H <sub>2</sub> 0, 300
calcium sulfate dihydrate	$caso_4 \cdot 2 H_2 O$	-1½ H <sub>2</sub> 0, 128		$-2 \text{ H}_2\text{ O}$ , 163
cobaltous nitrate hexahydrate	$Co(NO_3)_2 \cdot 6 H_2O$	<100		-3 H <sub>2</sub> 0, 50
ferrous chloride tetrahydrate	${ t FeC1}_2$			
sulfate heptahydrate	${\rm FeSO}_4^{-7}$ H <sub>2</sub> O	64; -6 H <sub>2</sub> 0, 100	(>20)	$-7 \text{ H}_2\text{O}, 300$
magnesium chloride hexahydrate	$MgC1_2 \cdot 6 H_2O$	d.116-8		ď.
manganous chloride tetrahydrate	$MnC1_2^{-4}$ $H_2^{0}$	58; -H <sub>2</sub> 0, 106		-4 H <sub>2</sub> 0, 198
sulfate monohydrate	$MnSO_4 \cdot H_2O$	stable 57-117		
potassium monohydrogen phosphate	$ ext{K}_2$ HPO $_4$	d.		
dihydrogen phosphate	$\mathrm{KH}_2\mathrm{PO}_4$	253	100	
bicarbonate	KHCO <sub>3</sub> .	d.100-200	(>0.02)	
sodium monohydrogen phosphate	${ m Na}_2{ m HPO}_4$			
dihydrogen phosphate mono- hydrate	$^{\mathrm{NaH_2PO_4}}\cdot^{\mathrm{H_2O}}$	-H <sub>2</sub> 0, 100		d.204
molybdate	$\text{Na}_2^{\text{MoO}_4} \cdot 2$ H <sub>2</sub> 0	-H <sub>2</sub> 0, 100		
bicarbonate	$_{\rm NaHCO_3}$	-co <sub>2</sub> , 270	11.6	
thiosulfate	Na <sub>2</sub> S <sub>2</sub> O <sub>3</sub>			
silicate nonahydrate	$^{\mathrm{Na}_{2}\mathrm{SiO}_{3}\cdot9}$ H $_{2}^{\mathrm{O}}$	40-48		-6 H <sub>2</sub> 0, 100

## APPENDIX D

# CONFIGURATION, COMPOSITION AND PREPARATION OF THE ALL-GLASS CONTAINER

## 1. CONFIGURATION AND COMPOSITION

The all-glass containers were LG 6770, 10-ml drying ampoules, made of Kimble KG-33 low expansion borosilicate glass, and were made by Lab Glass, Inc., Vineland, New Jersey.

## 1.1 PHYSICAL DESCRIPTION

The LG 6770 containers were round bottom drying ampoules with the following dimensions:

overall length	$184 \text{ mm} \pm 3 \text{ mm}$
stem length	95 mm
base length	89 mm
stem O.D.	$10 \text{ mm} \pm 0.5 \text{ mm}$
stem wall thickness	1.0 mm $\pm$ 0.2 mm
base O.D.	$15 \text{ mm} \pm 0.5 \text{ mm}$
base wall thickness	1.2 mm $\pm$ 0.2 mm

## 1.2 CHEMICAL COMPOSITION

The containers were made entirely of Kimble KG-33 glass (Owens-Illinois, Inc., Vineland, New Jersey). KG-33 glass is a low-expansion borosilicate glass.

#### 1.3 PHYSICAL PROPERTIES

The physical properties of KG-33 glass are:

strain point	515°C
annealing point	555°C
softening point	820°C
linear coef. of expansion	$32 \times 10^{-7} / ^{\circ} \text{C} (0-300 ^{\circ} \text{C})$
density (g/ml)	2.23
refractive index	1.47
transmission (2 mm)	92%
spec. heat (average for range 25°C to 175°C)	0.205 (ca1/g <sup>O</sup> C)
thermal cond. (20°C)	0.0028 (cal/sec cm °C)
Young's modulus psi	$8.9 \times 10^{6}$

## 1.4 BURSTING PRESSURE

Water was sealed into two containers according to the packaging procedure used for test substances except that the containers were held in liquid nitrogen during evacuation. The sealed containers were then placed in an aluminum-block tube heater and the temperature increased until they burst. By recording the temperature of the block it was possible to establish that the containers withstood pressures in excess of 700 psi before bursting.

## 1.5 U.S.P. TYPE

Ten containers, selected at random, were subjected to the U.S.P. Powdered Glass Test described in the <u>United States Pharmacopia</u>, seventeenth

revision, 1965, p.900. The test was carried out by Truesdail Laboratories, Inc., Los Angeles, California. Test results showed that glass used in the containers met requirements for U.S.P. Type I glass.

## 1.6 SINGLE BATCH CONTROL

All tubing used to construct the containers was Kimble K80200 standard wall glass tubing. Container stems were made of 10-mm tubing from Kimble lot number 124122866. Bases were made of 15-mm tubing from Kimble lot number 3121367. Kimble K80200 tubing is made of KG-33 glass.

## PREPARATION FOR STORAGE AND USAGE

## 2.1 PRELIMINARY WASH

The ampoules were inverted in stainless steel baskets, flushed with distilled water in the Heinicke model HW-5000E dishwasher (Heinicke Instruments Co., Hollywood, Florida) for two minutes, and then dried in an oven.

## 2.2 MARKING

Identification numbers were marked on the ampoules using a Glass Marker Grinder. The numbers were marked on the stem about 2 cm from the open end, and on the body about 3 cm below the shoulder of the ampoule.

## 2.3 WASHING AND ANNEALING

## 2.3.1 ACID WASH

The ampoules were immersed in A.C.S. grade, 90% Nitric Acid at about  $70^{\circ}$ C for 25-35 minutes.

## 2.3.2 FIRST RINSE

The ampoules were rinsed four times with purified water (U.S.P.). (This water was deionized water which met the U.S.P. test requirements.) The ampoules were filled completely and then drained on each rinse.

## 2.3.3 ANNEALING

The ampoules were annealed at  $560^{\circ}$ C for 15 minutes. The cooling rate was  $2^{\circ}$ C/min for the first  $150^{\circ}$ C.

## 2.3.4 FINAL RINSE

The ampoules were rinsed twice with purified water (U.S.P.), completely filled and then drained on each rinse. They were then rinsed twice more

with purified water (U.S.P.) which had been passed twice through Millipore Filter disks, pore size 0.22  $\mu\text{.}$  After the final rinse the ampoules were dried in an oven.

## 2.4 STORAGE

The ampoules were stored in a large, covered, polyethylene bucket until used.

## APPENDIX E

## THERMAL PROCESSING EQUIPMENT

#### 1. TUBE HEATERS

The thermal process environment was provided by two aluminum-block tube heaters. Each tube heater consisted of a heating base, an aluminum block, and a lid, made of Transite (an insulating material).

## 1.1 HEATING BASES

The heating bases were RSCo model 2127-A (Research Specialties Company, Richmond, California). They are heated electrically and rated by the manufacturer to cover the temperature range  $25-300^{\circ}\text{C} \pm 1^{\circ}\text{C}$ .

## 1.2 ALUMINUM BLOCKS

Each aluminum block had a capacity of forty-two, 15 mm tubes. The blocks were 11-5/8" long, 4-3/16" wide, and 3-1/2" deep. A sheet of Transite, 9/16" thick, was fastened to the top of each block. Forty-two holes, 5/8" in diameter, were drilled through the aluminum and Transite. Holes were 1 inch apart, measured center to center.

## 1.3 TEMPERATURE CONTROL

There were two principal sources of temperature variation in the heating blocks. At any one time a tube position near the center of the block could be warmer than one near the edge. Also, the temperature at a single tube position could vary over time. Calibration of the tube heaters (Section 2.2) showed that the difference in temperature between any two tube positions at one time was less than 2°C, and that the temperature at

any one tube position did not vary more than 1°C from its mean during the heat soak.

## 2. CALIBRATION OF EQUIPMENT

## 2.1 RECORDING POTENTIOMETER WITH TEMPERATURE SENSORS

A Foxboro model ERB12-30ML12-123 recording potentiometer (Foxboro Company, East Bridgewater, Massachusetts) and iron-constantan thermocouples were used to measure and monitor temperatures within the heating blocks. The recorder and thermocouples were calibrated against a National Bureau of Standards certified thermocouple by the Aeronutronic Division Standards Laboratory. Thermocouples were selected for similarity of voltage produced over the range of temperature measured in the investigation.

#### 2.2 TUBE HEATERS

At the start of each heat soak all specimens of a particular processing lot were placed in one of the tube heaters at one time. Because of a wide range of physical properties among the test substances, all specimens did not reach the specified temperature at the same time. The following calibration procedures were carried out to determine when to begin timing the 92-hour heat soak, and to determine if the difference in warmup times of specimens exceeded the  $\pm$  30 minutes allowed for variation in heat soak time. Stability and uniformity of temperature in the heating blocks were also determined.

## 2.2.1 WARMUP TIMES

The aluminum block and heating base were adjusted to a temperature near 135°C and allowed to equilibrate for 24 hours before final adjustment to the specified temperature was made. Iron-constantan thermocouples were placed in ten 15-mm test tubes. The tubes were filled with different test materials. Two contained 1 g sand, two contained 5 g sand, two were filled to a point five centimeters from the bottom with bentonite, two were filled to five centimeters with oil, and two were empty. These ten tubes were distributed in the heating block in such a manner that extremes of temperature would be indicated. Their temperatures were recorded for a 3-hour period. The entire set of tubes was then transferred to a test tube rack and cooled for one hour. This heating and cooling process was repeated five times. The mean warmup time (33 minutes) was used to determine the start of the 92-hour heat soak in the exposure of the test substances. The fastest warmup time was 18 minutes, the slowest was 52 minutes. The difference in warmup times was, therefore, within the range allowed for variation in heat soak time.

## 2.2.2 STABILITY AND UNIFORMITY OF TEMPERATURE

The aluminum block and heating base were adjusted to  $135^{\circ}\mathrm{C}$  and allowed 24 hours to equilibrate. Iron-constantan thermocouples were placed in ten, 15-mm test tubes containing about 5 g sand. These tubes were placed in representative holes in the heating block and their temperatures recorded for 48 hours. The temperature variation at any tube position was less than  $\pm$   $1^{\circ}\mathrm{C}$  during the 48 hours. The temperature at each tube position in the heating block was measured to determine the uniformity of temperature in the block. The steady-state temperatures were measured ten holes at a time. The arrays were so chosen that all holes were included in four trials. The difference between the highest and lowest temperatures in any group of ten positions was less than  $2^{\circ}\mathrm{C}$ .

## 3. MONITORING THE THERMAL PROCESS ENVIRONMENT

The temperatures of the heating blocks were monitored and recorded throughout each 92-hour heat soak period, including the associated warmup period.

# APPENDIX F

# TEST SEQUENCE

	Test No.	I Color Change	II Phase Change	III Loss of Volatiles	IV Particles in Solution	V pH Change	VI Differential Refractometry	VII X-Ray Diffraction Analysis	VIII Melting Point Determination	IX Barium Carbonate Pptn.	X Chromatography	XI Spectrophotometry	XII Fluorimetry	XIII Iodine Value Determination	XIV Amino Acid Analysis	XV Biological Assay	XVI Permanganate Reduction	XVII Colorimetry	XVIII Ferrocyanide Test
AMINO ACIDS																_			
8-Alanine L-Q-Alanine L-Arginine Hydrochloride L-Asparagine L-Aspartic Acid L-Cysteine Monohydrate Hydrochloride L-Cystine Dihydrochloride L-Glutamic Acid Hydrochloride L-Glutamine Glycine L-Histidine L-Hydroxyproline L-Isoleucine L-Leucine L-Leucine L-Phenylalanine L-Proline L-Serine L-Tryptophan L-Tyrosine L-Valine	de	1	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	3	4	5	6				7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7								
		1	2	J	4	J	U				7								
PROTEINS			_	•											,	, .			
Sodium Caseinate		1	2	3											4a	4Ъ			
MONOSACCHARIDES																			
Dextrose (α-D-Glucose) Levulose (β-D-Fructose) D-Galactose L-Arabinose D-Mannose		1 1	2 2 2 2 2	3 3	4 4 4	6 6	5 5 5				7 7 7 7								

		Color Change	Phase Change	Loss of Volatiles	Particles in Solution	pH Change	Differential Refractometry	X-Ray Diffraction Analysis	Melting Point Determination	Barium Carbonate Pptn.	Chromatography	Spectrophotometry	Fluorimetry	Iodine Value Determination	Amino Acid Analysis	Biological Assay	Permanganate Reduction	Colorimetry	Ferrocyanide Test
	Test No.	H	II	III	ΙΛ	Λ	ΙΛ	VII	VIII	ă	×	XI	XII	XIII	XIV	ΧV	XVI	XVII	XVIII
MONOSACCHARIDES, Contd.		_						·											
L-(+)Rhamnose Monohydrate N-Acetyl glucosamine  &Methyl-D-Glucoside D-Glucosamine Hydrochloride 2-Deoxy-D-Glucose Salicin D-Ribose		1 1 1 1 1 1	2 2 2 2 2 2 2 2	3 3 3 3 3 3	4 4 4 4 4 4	6 6 6 6 6 6	5 5 5 5 5 5				7 7 7 7 7 7								
OLIGOSACCHARIDES																			
D(-) Sucrose β-D-Maltose Monohydrate α-Lactose Monohydrate		1 1 1	2 2 2	3 3 3	4 4 4						5* 5* 5*	•							
POLYSACCHARIDES																			
Starch Inulin		1 1	2	3	4 4						5* 5*								
ALCOHOLS, POLYOLS																			
Ethanol Glycerin D-Mannitol Dulcitol (D-Galactitol)		1 1 1	2 2 2 2	3 3 3	4 4		4 4 7 7		5 5		5 5 6 6	6 6							
LIPIDS																			
Oleic Acid Sod. Salt Sodium Acetate Linoleic Acid		1 1 1	2 2 2	3 3 3	6	4	7 4				4 5	7		5 6			5		

<sup>\*</sup> after hydrolysis

		Color Change	Phase Change	Loss of Volatiles	Particles in Solution	pH Change	Differential Refractometry	X-Ray Diffraction Analysis	Melting Point Determination	Barium Carbonate Pptn.	Chromatography	Spectrophotometry	Fluorimetry	Iodine Value Determination	Amino Acid Analysis	Biological Assay	Permanganate Reduction	Colorimetry	Ferrocyanide Test	
	Test	· NO.	II	III	ΛI	٥	IA	VII	VIII	X	×	XI	XII	XIII	ΛΙΧ	XΛ	XVI	XVII	XVIII	
VITAMINS																				
Biotin Thiamine Hydrochloride Niacin (nicotinic acid) Riboflavin Pyridoxine Hydrochloride p-Aminobenzoic Acid i-Inositol (meso) Choline chloride Calcium Pantothenate Menadione Sodium Bisulfite Pimelic Acid		1 1 1 1 1 1 1 1 1 1	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	3 3 3 3 3 3 3 3 3 3	5 4 4 4 5 5 4 4 4 5		6 5 5 5 5 6 6 5 5 5 6		4 4 4		7 6 6 6 7 7 6 6 7	7	7 7 7					7 7		
PARTIAL HYDROLYSATES																				
Casitone Bacto-soytone Proteose Peptone #3		1 1 1	2 2 2	3 3 3	4 4 4						5 5 5				6a	61 61 61	)			
NUCLEIC ACID BASES																				
Adenine Guanine Hypoxanthine Xanthine Cytosine Orotic Acid Thymine Uracil		1 1 1 1 1 1 1	2	3 3 3	5 5 5		6 6 6 6 6 6				4 4 4 4 4 4 4 4	7 7 7 7 7 7 7								
BIOLOGICAL EXTRACTS																				
Yeast extract Beef extract Beef Heart for infusion Malt extract		1 1 1 1	2	3 3												4 4 4 4				

	Color Change	Phase Change	Loss of Volatiles	Particles in Solution	pH Change	Differential Refractometry	X-Ray Diffraction Analysis	Melting Point Determination	Barium Carbonate Pptn.	Chromatography	Spectrophotometry	Fluorimetry	Iodine Value Determination	Amino Acid Analysis	Biological Assay	Permanganate Reduction	Colorimetry	Ferrocyanide Test
Test	Ħ	II	III	IV	Λ	NI	VII	VIII	X	×	XI	XII	XIII	XIV	XX	XVI	XVII	XVIII
INORGANIC SALTS																		
Sodium Thiosulfate, Anhyd. Sodium Phosphate, Dibasic, Anhydrous	1	2 2	3	4	5	6	4											
Potassium Bicarbonate Manganous Sulfate, Monohydrate Ferrous Chloride	1	2 2 2	3 3 3	4 4	5 5	6 6			4									5
Ferrous Sulfate Ammonium Chloride Ammonium Molybdate Ammonium Phosphate, Dibasic	1 1 1 1	2 2 2 2	3 3 3	4	4 4	5	4											5
MISCELLANEOUS																		
Dipicolinic Acid Furfural Sodium Citrate, Dihydrate Succinic Acid Lactide Urea	1 1 1 1 1	2 2 2 2 2 2	3 3 3 3 3			4		4 4 4		4 5 4								
EXOBIOLOGICAL EXPERIMENT REAGENTS	_	_	J					7										
Sodium Formate Sodium Pyruvate L-α-Aspartyl-β-Naphthylamide L-Histidyl-β-Naphthylamide L-Phenylalanyl-β-Naphthylamide	1 1 1	2 2 2 2 2	3 3 3 3	4					5	6 5		4 4 4			5 5 5			

## APPENDIX G

## LABORATORY DATA

Laboratory data for all screening tests which have been completed are given in Tables I and II.

Candidate substances have been categorized into stability classes, based on results of tests which have been completed. Stability classes are:

- Class 1: Passed all tests.
- Class 2: Passed first three tests and failed one or more subsequent tests.
- Class 3: Failed the color test, because of a slight discoloration, and may have undergone only minimal damage (example: #41, Starch).
- Class 4: Failed the phase-change and loss of volatiles tests, but may have undergone only slight chemical change (example: #100, Urea).
- Class 5: Showed extensive decomposition.

Table I gives the stability class of each substance, indicates which tests have been completed, and reports the data from Tests I, II, and III. The data for Test I (Color Change) given in Munsell Color Values, are used to compare the color of heated and control specimens. The numbers represent Hue Value/Chroma. For example, 10 YR9/1 represents a color with Hue 10YR, Value 9 and Chroma 1. Chroma for neutral colors is zero and is not printed. The number N9.25/, therefore, represents neutral Hue, Value 9.25 and Chroma 0.

Descriptions of the phase of control and heated specimens are given as data for Test II (Phase Change). Under Test III (Loss of Volatiles), the weights of the heated specimens which were opened are given as sample weight. The weight changes reported are the differences in weight change between the control and heated specimen.

The data for Tests IV, VI, VII, VIII and IX are given in Table II. The appearance of solutions made from the heated and control specimens is reported under Test IV (Particles in Solution). Differential Refractometry (Test VI) data are given in terms of  $\Delta d$  and  $\Delta d'$ . These values are the differences between two numerical readings taken with the differential refractometer, and are proportional to the differences in the indexes of refraction. The  $\Delta d$  notation was the difference between the control and heated specimens and  $\Delta d'$  was the difference between the control and a 1% dilution of the control. Data for X-ray Diffraction Analysis (Test VII) are given as the percentages of candidate substances in the test sample. Samples were taken from control and heated specimens. Melting ranges of control and heated specimens are reported under Test VIII (Melting Point Determination). Melting ranges given are uncorrected. Data for Test IX (Barium Carbonate Precipitation) are in terms of the clarity of test solutions of control and heated specimens.

The results of all tests are reported as P or F for pass or fail. Test results which were marginal are enclosed in parentheses. The + symbol beside a P or F under test results indicates that a strong umpleasant odor was detected when the heated specimen was opened. The ++ symbol indicates that the odor was that of ammonia.

TABLE I

LABORATORY DATA FOR TESTS I, II AND III

										Test III	loss of	olatile
Substance and Quality	Stability	, Tests Completed	Code No.	Test I.	Color Change Heated Result	e Result	Control	Test II. Phase Change* Heated	Result	Sample wt.	Wt.Change mg Result	Resul
AMINO ACIDS							,		ı			1
8-Alanine, M.A.	5	1,11,111	1	/5.6N	N9.25/	ы	ff powd	wet cake		4.0499	?	Д.
L-o-Alanine, N.R.C.	1	I, II, III, IV, VI	2	/5.6N	/5.6N	Ь	ff cryst	cryst adhesion; not ff even after loosening; subl		4.0377	5.04	Δ, (
L-Arginine hydrochloride, N.R.C.	г	I, II, III, IV, VI	ю	/5.6N	N9.25/	ы	cryst	cryst subl		4.3411	5.5	<b>3.</b>
L-Asparagine, N.R.C.	5	1,11,111	4	/5.6N	10YR2/1	Œ	pmod	polym; some subl	Pari	4.5558	6.86-	Įz,
L-Aspartic acid, N.R.C.	1	I, II, III, IV, VI	2	/5.6N	N9.25/	Q.	cryst	cryst	д	4.2523	8. 0-	p.
L-Cysteine hydrochloride monohydrate, N.R.C.	5	1,11,111	9		ı	Ē	•	,	ĵε₄	5.0381	exploded	P4
L-Cystine dihydrochloride	en	1,11,111	7	N9.25/	10YR9/1	(F)	pwod	powd; some subl	д	4.0661	+3.7	<u>α</u> ,
L-Glutamic acid hydrochloride. C.P.	1	I,II,III,IV,VI	8	N9.25/	N9.25/	Δ.	coarse granl	coarse granl	д	5.0569	-0.5	Д
I_Clifamine N B C	· 50	III.III	6	/5.6N	5 78.5/2	Ŀ	pmod	fused cake	ĮΞι	2.9376	-0.2	Δ,
Contract N D	ır	I. II. III. IV. VI	10	/5.6N	5 Y 9 / 1	<u>[1</u> .	pmod	powd; liq on walls	д	3.9346	9.0+	A4
I_Wistidine N B C	, m	I.II.III.IV.VI	Π	N9.25/	5 79/1	(F)	loose cryst, not ff	ff cryst adhesion; no change in cryst shape	ы	3.3502	+0.3	Δ,
L-nistium, N.A.C.	) <del>-</del>	T TT TT TV VT	: :	NO 25/	/ 50 BN	<u>,</u>	ff gran1	cryst adhesion; ff after loosening	д	5,0419	-0.5	ρ.
4-Hydroxy-L-proline, N.K.C.	ન ત	I, II, III, IV, VI	7 2	NO 25.	/57 ON	4 P	ff cryst	ff cryst; some subl	д	1.5879	0	ρı
L-Isoleucine, N.R.C. (allo free)	7	1,11,111,10,V1	១ ;	/C7'6N	162.6N	ا بد	ff loafe	cryst adhesion;not ff after loosening;	Δ.	3.1312	+9.1	ρ,
L-Leucine (methionine free)	-	I,II,III,IV,VI	14	N9.25/	N9.25/	Δ,	71 TC413	no change in cryst shape	, Δ	3.4503	8 C	. Α
L-Lysine hydrochloride, N.R.C.	1	I,II,III,IV,VI	15	N9.25/	N9.25/	Д	granı	יייייייייייייייייייייייייייייייייייייי	, F	0200	) r	. 6
L-Methionine, N.R.C.	1	I, II, III, IV, VI	16	N9.5/	N9.25/	Д	pwod	powd; some subj	<u>1</u> 4 1	6076.7	0.7	ξ,
L-Phenylalanine, N.R.C.	1	I, II, III, IV, VI	17	/5.6N	/5.6N	д	ff granl	ff granl; some subl	ρ.,	2.949/	χ. Υ	<b>2</b> 4
L-Proline, N.R.C. (hydroxy-L-proline free)	5	1,11,111	18	/5.6N	5 Y9 /4	Į±,	pwod	fused cake & liq	Īz.	1.8623	-2.8	A,
L-Serine, N.R.C.	5	1,11,111	19		1	ы	•	1	ш	3.6414	exploded	P4
L-Thronning N R C (allo free)	e	I.II.III.IV.VI	20	N9.25/	5 Y9 / 1	(F)	pwod	pwod	ρι	3,9475	-0.3	щ
1-Tryntonhan N.R.C	2	I.II.III.VI	21	N9.25/	N9.25/	Д	pwod	pwod	Ь	1.3933	-0.7	д
Taypropries		T TT TT TV VT	33	/ 50 PM	/ 50 bN	Δ	pmod	pwod	(P)	1.476	4.0-	д
L-tyrosine, N.K.C.	٠,	1,11,111,11,11,1	2.5	/ S ON	/ 5 6N	, д	ff cryst	ff cryst; some subl (white & yellow)	(P)	2.2055	40.8	Δ,
L-Valine, N.K.C.	-1	1,11,111,11,1	7	(C: CH	(0.00	•						
PROTEINS	U	+	č	1/ 049	3/ 800.01	Þ	pood	caked powd	Ιŧι	1.2200	-8.7	£
Sodium caseinate (soluble casein)	n	1,11,111	<b>†</b> 7	1/616	o /out or	4	•					
CARBOHYDRATES												
Monosaccharides										,	;	1
N-Acety1-q-D-glucosamine	5	1,11,111	25	10YR9/1	10YR2/1	Į.,	pwod	polym	Ė	2.8114	-91.9	D-4
L-Arabinose, N.R.C.	5	1,11,111	27	N9.25/	10YR2/1	ĽΨ	pwod	polym	Ĺτι	4.2215	-39.9	<u>,</u> .
2-Deoxy-D-glucose, N.R.C.	5	1,11,111	28	N9.25/	10YR2/2	Γω	pwod	polym; some liq on walls	Ē	1.8971	-26.8	Pa <sub>1</sub>
Levulose (8-D-fructose), N.R.C.	5	1,11,111	29	/S.6N	10YR2/2	Ľι	ff gran1	polym; some liq on walls	Ŀ	4.2670	-201.3	P4
D-Galactose, N.R.C.	5	1,11,111	30	/5.6N	10 YR2 / 1	[24	pwod	ро1ут	Ŀ	2.9344	-13.3	Di.
D-Glucosamine hydrochloride	5	1,11,111	31		,	ĨΉ	:	•	[z.	4.1840	exploded	ſz,
Dextrose ( $\alpha$ -D-glucose), reagent, A.C.S.	5	1,11,111	32	/5.6N	10YR2/2	<u>14</u>	ff granl	polym	Ħ	5.0706	-22.8	
D-Mannose, N.R.C.	S	1,11,111	33	N9.25/	10YR2/1	īri	powd	polym	Ŀ	2.0622	-68.8	
α-Methy1-D-glucoside	3	I,II,III	34	/5.6N	5 Y 9/1	ſĿ,	pmod	pwod	<u>ρ</u>	2,6495	-2.2 g	
L(+)Rhamnose monohydrate, M.A.	5	I,II,III	35	/5.6N	5Y2/2	ſΣι	pmod	polym layer & liq layer	<u>124</u>	5.0088		۲
D-Salicin (Bact.), Puriss	1	I,II,III	36	/5.6N	N9.25/	ď	pwod	powd; some subl	ρ,	1.5374	N	
D-Ribose, N.R.C.	2	1,11,111	37	/5.6N	10YR2/1	Į±,	pood	polym	P4	1.9664	•;	(z)
Oligosaccharides											26.3	
lpha-Lactose monohydrate, U.S.P.	5	1,11,111	38	N9.25/	2.5 ¥4.5/4	ĵ±,	pmod	polym	Œ	4.5649	4.85	
8-D-Maltose monohydrate, N.R.C.	5	1,11,111	39	/S.6N	10 YR 2 / 2	F	pwod	polym	ш	1.8067	1	

of+18udeoge, Ipagent, A.C.S.	ž	Ι, ΙΙ, ΙΙΙ	07	N9.25/	5YR2/1	P <sup>2</sup>	large granl	polym	F 4	4.9581	-193.0	<u>A</u>
Polysaccharides												
Starch, soluble powder reagent, A.C.S.	6.0	1,11,111	14	N9.25/	2.5Y8.5/2	ţzı	pmod	bwo a			1	D*
fron 1 tn	2	1,11,111	77	/5.6N	10YR2/2	Ē	Dowd			3.2827	7.9	A
ALCOHOLS AND POLYOLS								FOLYE	F 2.	2.9657 -	-146.1	Œ
Dulcitol (D-galactitol), N.R.C.	3	I,II,III,VIII	43	/5.6N	5 Y 9/1	Ē	, paod					
Ethyl alcohol, pure (ethanol), U.S.P.	1	1,11,111	77	colorless	colorless	۵	140	caked powd; some decomp on walls	(P) 4.	4.6114	-0.7	a,
Glycerine (glycerol), reagent, A.C.S.	н	I,II,III	45		colorless	. Δ	114	btt	P 4.	4.7475	8.0-	d.
D-Mannitol, N.R.C.	1	I,II,III,VIII	46		000 25 / PN	, 0	110	liq	P 6.	6.7788	+3.2	Δ,
LIPIDS AND RELATED SUBSTANCES					107.00	4	pood	caked powd	P 2.	2.9350	-0.3	Δ,
Sodium acetate, reagent	1	I,II,III	47	/5.6N	/5.6N	p.	+ 0 2 2 2					
Linoleic acid, Puriss (99%)	e	1,11,111	84		5Y8.5/10	, E.	C1 yst.	cryst		4.8153	+2.9	Δ,
Oleic acid sodium salt (sodium oleate), practical	1	1,11,111	20	7,	2.5 29 /4	ę.	f1	11q E1			+0.3	<u>ρ</u>
YITAMINS								1	Y 7.8	2.8346	4.2	Δι
Para-aminobenzoic acid, U.S.P.	5	1,11,111	51		ı	Ç	;	!				
Biotin, crystalline	ı	I,II,III,VIII	52	N9.25/	N9.25/	, д,	cryst	CTUST	F 6	3.2237 exp	exploded 0 5	Er f
Choline chloride, crystalline (99%)	1	I,II,III	53		/5.6N	Δ.	large oran1				. · ·	4
1-Inositol (meso)	<b>-</b> -1	I,II,III	55		85.25	, д.	cryst	cryst			». o	e4 t
Niacin (nicotinic acid), U.S.P.	4	I,II,III	56		757 6N		- Poord	in a standard bank in a standard			0.0	л _
Calcium-D-pantorhenate, U.S.P.	۳	1.11.111	5.7			, (£	pour d	cased power, construction to sub.	_		5,0	<u>-</u> -
Pimelic acid. C.P.	'n	T TT TT	α L				7 7 10 10	cared power, some IIq on walls			-1.9	Ь
Duridowine monchiderochlowide (Witemin D6)	. ~	111111	2			<b>u</b> [	pwod.	rused cake	F 3.0	3.0229	-0.5	ъ ъ
Directic motoriyar oction to but	۰ -	111,111	٠,			_	powd	pwod	P 2.7	2.7986	-0.2	ъ
Kiboriawin B2)	→ ,	1,11,111	09	/16	9		bowd	pwod	P 1.3	1,3441	-3.6	4
Thiamine hydrochloride, (Vitamin B1), U.S.P.	m	1,11,111	61			ഥ	powd	powd; some subl	P 1.2	1.2976	+2.3	4
Menadione sodium bisulfite (water soluble Vitamin K)	رد د	1,11,111	62	N9.25/	5GY2/1 (	(F)	powd	polym			6.6-	. ρ.
ARTIAL HYDROLYSATES												
Casitone vitamin free, dehydrated	5	1,11,111	63	5 29/1 1	10YR4/6	(tu	pwod	slush	F 1.9	1.9147	-37.3	ŧ
Proteose Peptone No. 3	5	1,11,111	65	2.5Y8.5/4 1	10YR2/2	ţ.	large cryst	polym			7 62-	. 12
	5	1,11,111	99	2.5Y8.5/4 1	10YR2/2	<u></u>	p <sub>wc</sub> d	polym		•	-101.6	. [2
UCLEIC ACID BASES											)	4
		1,11,111	69	N /5.6N	/5.6N	ъ.	pmod	powd; some subl	P 2.7	2.7798	-0.3	Д
e, N.R.C.	e	1,11,111	70	s /s.en	5 Y 9 / 1	F	pood	pmod			-2.5	, t
Guanine	7	1,11,111	71	N /5.6N	/5.6N	P 1	pmod	powd; some subl	P 2.9		0.1-	
	3	1,11,111	72	5Y8.5/1 5	5Y9/2	F	pwoci	pwod				, (a)
rboxylic acid)	m	1,11,111	73	N9.5 1	10YR9/1	<u> </u>	p.o.o.d	caked powd				, A
e (5-mothyluracil)	1	1,11,111	74	5 Y8.5/1 5	5Y8.5/1	P f	£1	f1			-2.0	, д
Uracil	1	1,11,111	75	5 1/615	5 49/1	7 1	pmod	pwod			6.0-	. 4
Xanthine		111 11	3.5	NO 5./	9.5/	F.	משטים	pwou	P 3.4893	1		- Δ
ATACTCAL EXTRACTS										1	ı	

	•						į					
			;	1,0000	1/ 5/11/01	£	secto (foams)	nasts (no fram): some subl	Д	4.7279	-59.3	[24
	7	I,II,III	1	IOYKZ/I	10 IR2 / 1	4	paste (toamy)					i
	u	III.III	78	2.578/2	10YR7/4	Ľω	pmod	caked powd	ц	1.4905	-25.6	ŧ,
deart for incusion	٠ ١٠	I,II,III	42	2.579/2	10YR2/2	<u> </u>	pood	polym	Ēz,	2,9503		ĬΨ
Mair extract	1						•	•	T.	1 0000	2 70	ţ.
Yeast extract	5	I,II,III	80	10YR9/2	10YR2/1	T24	pwod	s lush	24	1.9906	0.00	4
INORGANIC SALITS								:	í			F
Ammonium chloride, granular, reagent, A.C.S.	1	1,11,111	81	N9.5/	/5.6N	<b>d</b>	granl	granl; some subl	14	5.0215	0.1.	<b>L,</b> ,
Ammonium molubdate crustal, reagent, A.C.S.	5	1,11,111	82	10Y9/1	7.5YR9/2	(F)	loose granl	solid cake; some 11q but reabsorbed	Įz.	5.0857	-1.8	Д
Authorite and a second	-	T. TT. TT. VII	83	N9.5/	N9.25/	Д	ff granl	gran1 adhesion	Δ,	4.9048	-3.9	д
Ammonium phosphare, dibasic, crystar, reagent, a.c.o.	٠,	±, ±=, ±==, ===	΄ /α	5 V8 5 1/4	2/ 5007 /4	(F)	granl	solid cake; some liq	ы	2.5397	-2.8	Ы
Ferrous chloride, crystal, reagent	n	1,1,1,1	; ;			) f	1	l uero	д	3,4890	+3.0	Д
Ferrous sulfate, crystal, reagent A.C.S.	-	I,II,III	85	//N	/ X	٠,	granı		p	6 0577	70	Þ
Manganous sulfate monohydrate, powder, reagent, A.C.S.	1	1,11,111	98	/5.6N	/5.6N	<u>а</u>	pwod	powd	4	1100.0	7.01	4 1
Dotassium hicarbonate crystal, reagent, A.C.S.	1	I,II,III,IV,IX	87	/5.6N	/5.6N	ы	cryst	cryst	щ	5.0438	6.0-	بد
Gallian - Leaster ditario reasont & C	-	I.II.III.VII	88	79.5V	/5.6N	ρ.,	pmod	pwod	д	4.9839	-0.1	գ
Sodium phosphare, albasic, reagent, a.c.o.		III,III	68	/5.6N	/5.6N	ρ.,	granl	granl	ρ.	5.0647	+0.3	e,
Southum Chilosofface, teaberin	,											
MISCELLANEOUS								•	p	3177 1	,	Δ
Dipicolinic acid (2,6-pyridine dicarboxylic acid)	1	1,11,111	90	N9.25/	N9.25/	ы	pmod	powd	4 9	01//1	2	•
Furfural, reagent	2	1,11	91	5Y9/2	10YR2/2	ш	liq	119	<b>2</b> 4 1	0.5265		. ,
Lactide, reagent	3	1,11,111	92	10 YR9 /2	2.5YR3/4	Ŀ	granl	gran1 adhesion	щ	1.9824	?: 	<b>.</b>
Sodium citrate dihydrate, crystal, reagent	2	I,II	93	N9.25/	N9.25/	Ь	ff granl	solid cake;liq on walls;reabsorbed 11q & broke ampoule	Į.	5.0547	, ,	
Succinic acid, crystal, reagent		I,II,III,VIII	94	N9.25/	N9.25/	Ь	loose granl	granl adhesion	d,	4.9472	+1.4	щ
1 Accessive Company of the Compa	m	1,11,111	95	/5.6N	5 Y 9 / 1	Œ	pwod	pwod	Ь	0.1914	<b>5.0</b> -	A.
The state of the s	· Lr	111.111	96	5 79/1	2.5YR6/2	Ħ	fine powd	fused cake; some subl	Ēų	0.1760	40.5	Д
L-HISLIGHT-D-Haphen true, 11,55.	, L	_,, T TT TT	44	1/6/25	10YR3/6	ſz,	fine powd	fused cake	ы	0,1909	-1.9	Dz.
L-Phenylalanyl- $\beta$ -naphthylamide, M.A.	o ,	111 (11 (1 ) )	; ;	1 1 2 2	, , , , ,	۵	fine crust	cryst adhesion	д	4.8056	+1.3	д
Sodium formate, crystal, reagent	_	I, II, III, IV, LX	ž,	7C. 6N	10.6N	4	TIME CLYSE		í	0077	7 07	4

II,II,I I

I,II,III,IV,IX I,II,III I,II,III Urea, crystal, reagent, A.C.S.

Sodium pyruvate, reagent

Abbrevia	tions	nsed	to	describe	phase	οĘ	test	Abbreviations used to describe phase of test substances (Test III):	(Test	:(111	
££	free	free flowing	80		f1		£1	flakes			
powd	powder	er			subl	ŭ	sn	sublimate			
cryst	crystals	tals			liq	_	1;	liquid			
granl	granules	ules			polym	m/	М	polymer			

cryst granl

G-3

71.

, ‡

-59.3

2.4403

fused cake; some subl particles adhesion cryst adhesion

coarse granl

powd

10YR5/8 /5.6N

5.29/5

66 100

/5.6N

/5.6N

+1.3 7.65-

	Code	Test IV	. Particl	es in Sol	ution	Control & Heate
Substance and Quality	No.	g/ml H <sub>2</sub> O	Control	Heated	Result	% Conc.
AMINO ACIDS						
$L-\alpha$ -Alanine, N.R.C.	2	0.2500/3	clear	clear	P	7.70
L-Arginine hydrochloride, N.R.C.	3	1.500/3	clear	clear	P	33.33
L-Aspartic acid, N.R.C.	5	0.0575/30	clear	clear	P	0.19
L-Glutamic acid hydrochloride, C.P.	8	0.3750/3	clear	clear	P	11.12
Glycine, N.R.C.	10	0.7500/3	clear	clouded	F	20.00
L-Histidine, N.R.C.	11	0.6000/30	clear	clear	P	1.96
4-Hydroxy-L-proline, N.R.C.	12	0.7500/3	clear	clear	P	20.00
L-Isoleucine, N.R.C. (allo free)	13	0.3250/30	clear	clear	P	1.07
L-Leucine (methionine free)	14	0.0130/30	clear	clear	P	0.0433
L-Lysine hydrochloride, N.R.C.	15	1.5000/3	clear	clear	P	33.33
L-Methionine, N.R.C.	16	0.1000/3	clear	clear	P	3.23
L-Phenylalanine, N.R.C.	17	0.4500/30	clear	clear	P	1.48
L-Threonine, N.R.C. (allo free)	20	0.1500/3	clear	clear	P	4.76
L-Tryptophan, N.R.C.	21	0.1500/30	clear	clear	P	0.498
L-Tyrosine, N.R.C.	22	0.0100/ <b>3</b> 0	clear	clear	P	0.0333
L-Valine, N.R.C.	23	0.1250/3	clear	clear	P	4.0
ALCOHOLS AND POLYOLS						
Dulcitol (D-galactitol), N.R.C.	43					
D-Mannitol, N.R.C.	46					
VITAMINS						
Biotin, crystalline	52					
INORGANIC SALTS						
Ammonium phosphate, dibasic, crystal, reagent, A.C.S.	83					
Potassium bicarbonate, crystal, reagent, A.C.S.	87	1.0/3	clear	clear	P	
Sodium phosphate, dibasic, reagent, A.C.S.	88					
MISCELLANEOUS						
Succinic acid, crystal, reagent	94					
Sodium formate, crystal, reagent	98	1.0/2	clear	clear	P	

TABLE II

LABORATORY DATA FOR TESTS IV, VI, VII, VIII AND IX

Test VI. Differential Refractometry				•		<i>:</i>		
, Same Conc.	Two Control	s, 1% Diff.	in Conc		Test VII. X-r	ay Diffract	ion Analysis	Tes
∆d	% Conc.	% Conc.	∆d '	Result	Control	Heated	Result	
	·		· · · · · · · · · · · · · · · · · · ·	· · · · · · · · · · · · · · · · · · ·			<del></del>	
10	7 70	7 (1	001	_				
18	7.70	7.61	231	P				
82	33.33	33.0	668	P				
10	0.19	0.188	61	P				
15	11.12	11.00	239	P				
1773	20.00	19.8	1335	F				
25	1.96	1.94	65	P				
31	20.00	1.98	443	P				
91	1.07	1.06	34	F				
14	0.0433	0.0429	43	P				
149	33.33	33.00	379	P				
20	3.23	3.20	33	P				
15	1.48	1.46	88	P				
41	4.76	4.71	67	P				
11	0.498	0.493	11	<b>(</b> F)				
21	0.0333	0.0330	43	P				
47	4.0	3.96	61	P				

25%	25%	P
100%	100%	п

t VIII. Melting Point Determination, °C				TEST IN	Dallu		Precipitation	Code
Control	Heated	Mixture	Result	Co	ntrol_	Heated	Result	No.
								2
								2
								3
								5
								8
								10
								11
								12
								13
								14
								15
								16
								17
								20
								21
								22
								23
186-188	186-188	186-188	P					43
165-167	165-167	165-167	P					46
000 000	222 222		_					52
229-230	229-230	229-230	P					34
								83
					lear	clear	P	.8
					LCUL	Clear		8
								0
187-189	187-189	187-189	P					9

clear clear P 98